Introduction to finite element methods

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The finite element method is a powerful tool for solving differential ec The method can easily deal with complex geometries and higher-order applications of the solution. Figure 1 shows a two-dimensional domain with a not geometry. The idea is to divide the domain into triangles (elements) a a polynomial approximations to the unknown functions on each triang method glues these piecewise approximations together to find a global standard quadratic polynomials over the triangles are particularly po

Many successful numerical methods for differential equations, including finite element method, aim at approximating the unknown function by

$$u(x) = \sum_{i=0}^{N} c_i \psi_i(x),$$

where $\psi_i(x)$ are prescribed functions and c_0, \ldots, c_N are unknown coeffices be determined. Solution methods for differential equations utilizing (have a principle for constructing N+1 equations to determine c_0, \ldots, c_I there is a machinery regarding the actual constructions of the equat c_0, \ldots, c_N , in a particular problem. Finally, there is a solve phase for conthe solution c_0, \ldots, c_N of the N+1 equations.

Especially in the finite element method, the machinery for construction discrete equations to be implemented on a computer is quite comprehension many mathematical and implementational details entering the scene at t

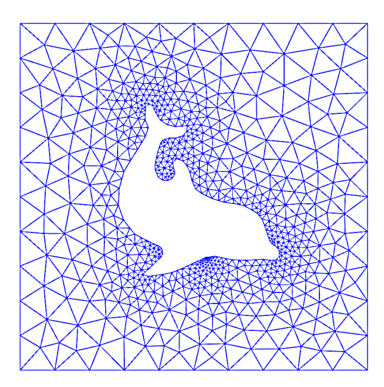


Figure 1: Domain for flow around a dolphin.

me. From an ease-of-learning perspective it can therefore be wise to follow n idea of Larson and Bengzon [1] and introduce the computational machinery or a trivial equation: u=f. Solving this equation with f given and u on the rm (1) means that we seek an approximation u to f. This approximation roblem has the advantage of introducing most of the finite element toolbox, ut with postponing demanding topics related to differential equations (e.g., itegration by parts, boundary conditions, and coordinate mappings). This is ne reason why we shall first become familiar with finite element approximation efore addressing finite element methods for differential equations.

First, we refresh some linear algebra concepts about approximating vectors vector spaces. Second, we extend these concepts to approximating functions function spaces, using the same principles and the same notation. We present tamples on approximating functions by global basis functions with support aroughout the entire domain. Third, we introduce the finite element type of scal basis functions and explain the computational algorithms for working with ach functions. Three types of approximation principles are covered: 1) the least quares method, 2) the L_2 projection or Galerkin method, and 3) interpolation r collocation.

1 Approximation of vectors

We shall start with introducing two fundamental methods for determing coefficients c_i in (1) and illustrate the methods on approximation of because vectors in vector spaces give a more intuitive understanding than directly with approximation of functions in function spaces. The expression functions will be trivial as soon as the fundamental in understood.

The first method of approximation is called the *least squares method* consists in finding c_i such that the difference u-f, measured in some minimized. That is, we aim at finding the best approximation u to f (norm). The second method is not as intuitive: we find u such that the u-f is orthogonal to the space where we seek u. This is known as pr or we may also call it a *Galerkin method*. When approximating vect functions, the two methods are equivalent, but this is no longer the case applying the principles to differential equations.

1.1 Approximation of planar vectors

Suppose we have given a vector $\mathbf{f} = (3, 5)$ in the xy plane and that we approximate this vector by a vector aligned in the direction of the vector Figure 2 depicts the situation.

We introduce the vector space V spanned by the vector $\psi_0 = (a, b)$:

$$V = \operatorname{span} \{ \boldsymbol{\psi}_0 \} .$$

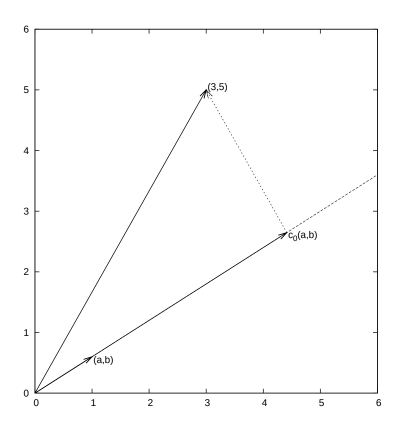
We say that ψ_0 is a basis vector in the space V. Our aim is to fivector $\mathbf{u} = c_0 \psi_0 \in V$ which best approximates the given vector $\mathbf{f} = (\mathbf{r})$ reasonable criterion for a best approximation could be to minimize the left the difference between the approximate \mathbf{u} and the given \mathbf{f} . The difference $\mathbf{e} = \mathbf{f} - \mathbf{u}$, has its length given by the norm

$$||e|| = (e, e)^{\frac{1}{2}},$$

where (e, e) is the *inner product* of e and itself. The inner product, als scalar product or dot product, of two vectors $\mathbf{u} = (u_0, u_1)$ and $\mathbf{v} = (u_0, u_1)$ and

$$(\boldsymbol{u},\boldsymbol{v})=u_0v_0+u_1v_1.$$

Remark 1. We should point out that we use the notation (\cdot, \cdot) for two things: (a, b) for scalar quantities a and b means the vector starting in the and ending in the point (a, b), while (u, v) with vectors u and v means the product of these vectors. Since vectors are here written in boldface for should be no confusion. We may add that the norm associated with the product is the usual Eucledian length of a vector.



igure 2: Approximation of a two-dimensional vector by a one-dimensional ector.

temark 2. It might be wise to refresh some basic linear algebra by consulting textbook. Exercises 1 and 2 suggest specific tasks to regain familiarity with indamental operations on inner product vector spaces.

'he least squares method. We now want to find c_0 such that it minimizes e||. The algebra is simplified if we minimize the square of the norm, $||e||^2 = c$, e), instead of the norm itself. Define the function

$$E(c_0) = (\mathbf{e}, \mathbf{e}) = (\mathbf{f} - c_0 \psi_0, \mathbf{f} - c_0 \psi_0). \tag{4}$$

/e can rewrite the expressions of the right-hand side in a more convenient form or further work:

$$E(c_0) = (\mathbf{f}, \mathbf{f}) - 2c_0(\mathbf{f}, \psi_0) + c_0^2(\psi_0, \psi_0).$$
 (5)

he rewrite results from using the following fundamental rules for inner product paces:

$$(\alpha \boldsymbol{u}, \boldsymbol{v}) = \alpha(\boldsymbol{u}, \boldsymbol{v}), \quad \alpha \in \mathbb{R},$$

$$(u + v, w) = (u, w) + (v, w),$$

$$(\boldsymbol{u}, \boldsymbol{v}) = (\boldsymbol{v}, \boldsymbol{u}).$$

Minimizing $E(c_0)$ implies finding c_0 such that

$$\frac{\partial E}{\partial c_0} = 0.$$

Differentiating (5) with respect to c_0 gives

$$\frac{\partial E}{\partial c_0} = -2(\boldsymbol{f}, \boldsymbol{\psi}_0) + 2c_0(\boldsymbol{\psi}_0, \boldsymbol{\psi}_0).$$

Setting the above expression equal to zero and solving for c_0 gives

$$c_0 = rac{(oldsymbol{f}, oldsymbol{\psi}_0)}{(oldsymbol{\psi}_0, oldsymbol{\psi}_0)},$$

which in the present case with $\psi_0 = (a, b)$ results in

$$c_0 = \frac{3a + 5b}{a^2 + b^2} \,.$$

For later, it is worth mentioning that setting the key equation (9) can be rewritten as

$$(\boldsymbol{f} - c0\boldsymbol{\psi}_0, \boldsymbol{\psi}_0) = 0.$$

or

$$(\boldsymbol{e},\boldsymbol{\psi}_0)=0$$
.

The projection method. We shall now show that minimizing $||e||^2$ that e is orthogonal to any vector v in the space V. This result is visual clear from Figure 2 (think of other vectors along the line (a, b): all of the lead to a larger distance between the approximation and f). To see the mathematically, we express any $v \in V$ as $v = s\psi_0$ for any scalar parameter that two vectors are orthogonal when their inner product vanish

alculate the inner product

$$\begin{split} (\boldsymbol{e}, s\psi_0) &= (\boldsymbol{f} - c_0 \psi_0, s\psi_0) \\ &= (\boldsymbol{f}, s\psi_0) - (c_0 \psi_0, s\psi_0) \\ &= s(\boldsymbol{f}, \psi_0) - sc_0(\psi_0, \psi_0) \\ &= s(\boldsymbol{f}, \psi_0) - s\frac{(\boldsymbol{f}, \psi_0)}{(\psi_0, \psi_0)} (\psi_0, \psi_0) \\ &= s\left((\boldsymbol{f}, \psi_0) - (\boldsymbol{f}, \psi_0)\right) \\ &= 0. \end{split}$$

herefore, instead of minimizing the square of the norm, we could demand that is orthogonal to any vector in V. This method is known as *projection*, because is the same as projecting the vector onto the subspace. (The approach can so be referred to as a Galerkin method as explained at the end of Section 1.2.) Mathematically the projection method is stated by the equation

$$(\boldsymbol{e}, \boldsymbol{v}) = 0, \quad \forall \boldsymbol{v} \in V.$$
 (13)

n arbitrary $v \in V$ can be expressed as $s\psi_0$, $s \in \mathbb{R}$, and therefore (13) implies

$$(\boldsymbol{e}, s\boldsymbol{\psi}_0) = s(\boldsymbol{e}, \boldsymbol{\psi}_0) = 0,$$

hich means that the error must be orthogonal to the basis vector in the space :

$$(e, \psi_0) = 0$$
 or $(f - c_0 \psi_0, \psi_0) = 0$.

he latter equation gives (10) and it also arose from least squares computations ι (12).

.2 Approximation of general vectors

et us generalize the vector approximation from the previous section to vectors 1 spaces with arbitrary dimension. Given some vector f, we want to find the est approximation to this vector in the space

$$V = \operatorname{span} \{ \psi_0, \dots, \psi_N \}$$
.

We assume that the basis vectors ψ_0, \ldots, ψ_N are linearly independent so that one of them are redundant and the space has dimension N+1. Any vector $\in V$ can be written as a linear combination of the basis vectors,

$$\boldsymbol{u} = \sum_{j=0}^{N} c_j \boldsymbol{\psi}_j,$$

here $c_j \in \mathbb{R}$ are scalar coefficients to be determined.

The least squares method. Now we want to find c_0, \ldots, c_N , such t the best approximation to f in the sense that the distance (error) e = is minimized. Again, we define the squared distance as a function of parameters c_0, \ldots, c_N ,

$$E(c_0, \dots, c_N) = (\boldsymbol{e}, \boldsymbol{e}) = (\boldsymbol{f} - \sum_j c_j \boldsymbol{\psi}_j, \boldsymbol{f} - \sum_j c_j \boldsymbol{\psi}_j)$$
$$= (\boldsymbol{f}, \boldsymbol{f}) - 2 \sum_{j=0}^N c_j (\boldsymbol{f}, \boldsymbol{\psi}_j) + \sum_{p=0}^N \sum_{q=0}^N c_p c_q (\boldsymbol{\psi}_p, \boldsymbol{\psi}_q).$$

Minimizing this E with respect to the independent variables c_0, \ldots, c_N is c by requiring

$$\frac{\partial E}{\partial c_i} = 0, \quad i = 0, \dots, N.$$

The second term in (14) is differentiated as follows:

$$rac{\partial}{\partial c_i} \sum_{j=0}^N c_j(oldsymbol{f}, oldsymbol{\psi}_j) = (oldsymbol{f}, oldsymbol{\psi}_i),$$

since the expression to be differentiated is a sum and only one term, c contains c_i and this term is linear in c_i . To understand this differential detail, write out the sum specifically for, e.g. N=3 and i=1.

The last term in (14) is more tedious to differentiate. We start with

$$\frac{\partial}{\partial c_i} c_p c_q = \begin{cases} 0, & \text{if } p \neq i \text{ and } q \neq i, \\ c_q, & \text{if } p = i \text{ and } q \neq i, \\ c_p, & \text{if } p \neq i \text{ and } q = i, \\ 2c_i, & \text{if } p = q = i, \end{cases}$$

Then

$$rac{\partial}{\partial c_i}\sum_{p=0}^N\sum_{q=0}^N c_p c_q(oldsymbol{\psi}_p,oldsymbol{\psi}_q) = \sum_{p=0,p
eq i}^N c_p(oldsymbol{\psi}_p,oldsymbol{\psi}_i) + \sum_{q=0,q
eq i}^N c_q(oldsymbol{\psi}_q,oldsymbol{\psi}_i) + 2c_i(oldsymbol{\psi}_q,oldsymbol{\psi}_q,oldsymbol{\psi}_q,oldsymbol{\psi}_q)$$

The last term can be included in the other two sums, resulting in

$$rac{\partial}{\partial c_i} \sum_{p=0}^N \sum_{q=0}^N c_p c_q(\boldsymbol{\psi}_p, \boldsymbol{\psi}_q) = 2 \sum_{j=0}^N c_i(\boldsymbol{\psi}_j, \boldsymbol{\psi}_i) \,.$$

It then follows that setting

$$\frac{\partial E}{\partial c_i} = 0, \quad i = 0, \dots, N,$$

ads to a linear system for c_0, \ldots, c_N :

$$\sum_{j=0}^{N} A_{i,j} c_j = b_i, \quad i = 0, \dots, N,$$
(18)

here

$$A_{i,j} = (\psi_i, \psi_j), \tag{19}$$

$$b_i = (\boldsymbol{\psi}_i, \boldsymbol{f}). \tag{20}$$

/e have changed the order of the two vectors in the inner product according to ..1):

$$A_{i,j} = (\boldsymbol{\psi}_i, \boldsymbol{\psi}_i) = (\boldsymbol{\psi}_i, \boldsymbol{\psi}_j),$$

mply because the sequence i-j looks more aesthetic.

'he Galerkin or projection method. In analogy with the "one-dimensional" cample in Section 1.1, it holds also here in the general case that minimizing the istance (error) e is equivalent to demanding that e is orthogonal to all $v \in V$:

$$(\boldsymbol{e}, \boldsymbol{v}) = 0, \quad \forall \boldsymbol{v} \in V.$$
 (21)

ince any $v \in V$ can be written as $v = \sum_{i=0}^{N} c_i \psi_i$, the statement (21) is quivalent to saying that

$$(oldsymbol{e}, \sum_{i=0}^N c_i oldsymbol{\psi}_i) = 0,$$

or any choice of coefficients c_0, \ldots, c_N . The latter equation can be rewritten as

$$\sum_{i=0}^N c_i(oldsymbol{e},oldsymbol{\psi}_i) = 0$$
 .

this is to hold for arbitrary values of c_0, \ldots, c_N we must require that each rm in the sum vanishes,

$$(e, \psi_i) = 0, \quad i = 0, \dots, N.$$
 (22)

hese N+1 equations result in the same linear system as (18):

$$(oldsymbol{f} - \sum_{j=0}^N c_j oldsymbol{\psi}_j, oldsymbol{\psi}_i) = (oldsymbol{f}, oldsymbol{\psi}_i) - \sum_{j \in \mathcal{I}_s} (oldsymbol{\psi}_i, oldsymbol{\psi}_j) c_j = 0,$$

nd hence

$$\sum_{j=0}^N (oldsymbol{\psi}_i, oldsymbol{\psi}_j) c_j = (oldsymbol{f}, oldsymbol{\psi}_i), \quad i = 0, \dots, N\,.$$

So, instead of differentiating the $E(c_0, \ldots, c_N)$ function, we could sim (21) as the principle for determining c_0, \ldots, c_N , resulting in the N+1 et (22).

The names least squares method or least squares approximation are since the calculations consists of minimizing $||e||^2$, and $||e||^2$ is a sum of of differences between the components in f and u. We find u such the sum of squares is minimized.

The principle (21), or the equivalent form (22), is known as pr Almost the same mathematical idea was used by the Russian mathen Boris Galerkin¹ to solve differential equations, resulting in what is widely as Galerkin's method.

2 Approximation of functions

Let V be a function space spanned by a set of basis functions ψ_0, \ldots, ψ

$$V = \operatorname{span} \{\psi_0, \dots, \psi_N\},\$$

such that any function $u \in V$ can be written as a linear combination of t functions:

$$u = \sum_{j \in \mathcal{I}_s} c_j \psi_j \,.$$

The index set \mathcal{I}_s is defined as $\mathcal{I}_s = \{0, \dots, N\}$ and is used both for contation and for flexibility in the numbering of elements in sequences.

For now, in this introduction, we shall look at functions of a single x: u = u(x), $\psi_i = \psi_i(x)$, $i \in \mathcal{I}_s$. Later, we will almost trivially ext mathematical details to functions of two- or three-dimensional physica. The approximation (23) is typically used to discretize a problem in space methods, most notably finite differences, are common for time discretal although the form (23) can be used in time as well.

2.1 The least squares method

Given a function f(x), how can we determine its best approximation u(A) natural starting point is to apply the same reasoning as we did for in Section 1.2. That is, we minimize the distance between u and f. It this requires a norm for measuring distances, and a norm is most convidefined through an inner product. Viewing a function as a vector of it many point values, one for each value of x, the inner product could int

¹http://en.wikipedia.org/wiki/Boris_Galerkin

e defined as the usual summation of pairwise components, with summation eplaced by integration:

$$(f,g) = \int f(x)g(x) dx$$
.

o fix the integration domain, we let f(x) and $\psi_i(x)$ be defined for a domain $\subset \mathbb{R}$. The inner product of two functions f(x) and g(x) is then

$$(f,g) = \int_{\Omega} f(x)g(x) \, dx. \qquad (24)$$

The distance between f and any function $u \in V$ is simply f - u, and the quared norm of this distance is

$$E = (f(x) - \sum_{j \in \mathcal{I}_s} c_j \psi_j(x), f(x) - \sum_{j \in \mathcal{I}_s} c_j \psi_j(x)).$$
 (25)

ote the analogy with (14): the given function f plays the role of the given ector f, and the basis function ψ_i plays the role of the basis vector ψ_i . We can ewrite (25), through similar steps as used for the result (14), leading to

$$E(c_i, \dots, c_N) = (f, f) - 2 \sum_{j \in \mathcal{I}_s} c_j(f, \psi_i) + \sum_{p \in \mathcal{I}_s} \sum_{q \in \mathcal{I}_s} c_p c_q(\psi_p, \psi_q).$$
 (26)

linimizing this function of N+1 scalar variables $\{c_i\}_{i\in\mathcal{I}_s}$, requires differentiation ith respect to c_i , for all $i\in\mathcal{I}_s$. The resulting equations are very similar to rose we had in the vector case, and we hence end up with a linear system of reform (18), with basically the same expressions:

$$A_{i,j} = (\psi_i, \psi_j), \tag{27}$$

$$b_i = (f, \psi_i). (28)$$

.2 The projection (or Galerkin) method

s in Section 1.2, the minimization of (e, e) is equivalent to

$$(e, v) = 0, \quad \forall v \in V. \tag{29}$$

his is known as a projection of a function f onto the subspace V. We may also all it a Galerkin method for approximating functions. Using the same reasoning f in (21)-(22), it follows that (29) is equivalent to

$$(e, \psi_i) = 0, \quad i \in \mathcal{I}_s. \tag{30}$$

iserting e = f - u in this equation and ordering terms, as in the multiimensional vector case, we end up with a linear system with a coefficient matrix '7) and right-hand side vector (28). Whether we work with vectors in the plane, general vectors, or fu in function spaces, the least squares principle and the projection or C method are equivalent.

2.3 Example: linear approximation

Let us apply the theory in the previous section to a simple problem: parabola $f(x) = 10(x-1)^2 - 1$ for $x \in \Omega = [1, 2]$, find the best approx u(x) in the space of all linear functions:

$$V = \operatorname{span} \{1, x\}.$$

With our notation, $\psi_0(x) = 1$, $\psi_1(x) = x$, and N = 1. We seek

$$u = c_0 \psi_0(x) + c_1 \psi_1(x) = c_0 + c_1 x,$$

where c_0 and c_1 are found by solving a 2×2 the linear system. The comatrix has elements

$$A_{0,0} = (\psi_0, \psi_0) = \int_1^2 1 \cdot 1 \, dx = 1,$$

$$A_{0,1} = (\psi_0, \psi_1) = \int_1^2 1 \cdot x \, dx = 3/2,$$

$$A_{1,0} = A_{0,1} = 3/2,$$

$$A_{1,1} = (\psi_1, \psi_1) = \int_1^2 x \cdot x \, dx = 7/3.$$

The corresponding right-hand side is

$$b_1 = (f, \psi_0) = \int_1^2 (10(x-1)^2 - 1) \cdot 1 \, dx = 7/3,$$

$$b_2 = (f, \psi_1) = \int_1^2 (10(x-1)^2 - 1) \cdot x \, dx = 13/3.$$

Solving the linear system results in

$$c_0 = -38/3, \quad c_1 = 10,$$

and consequently

$$u(x) = 10x - \frac{38}{3}$$
.

Figure 3 displays the parabola and its best approximation in the space linear functions.

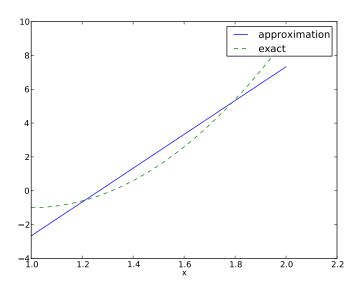


Figure 3: Best approximation of a parabola by a straight line.

.4 Implementation of the least squares method

ymbolic integration. The linear system can be computed either symbolially or numerically (a numerical integration rule is needed in the latter case). Lere is a function for symbolic computation of the linear system, where f(x) is liven as a sympy expression f involving the symbol x, psi is a list of expressions or $\{\psi_i\}_{i\in\mathcal{I}_x}$, and Omega is a 2-tuple/list holding the limits of the domain Ω :

```
import sympy as sp
lef least_squares(f, psi, Omega):
   N = len(psi) - 1
   A = sp.zeros((N+1, N+1))
   b = sp.zeros((N+1, 1))
   x = sp.Symbol('x')
   for i in range(N+1):
       for j in range(i, N+1):
           A[i,j] = sp.integrate(psi[i]*psi[j],
                                  (x, Omega[0], Omega[1]))
           A[j,i] = A[i,j]
       b[i,0] = sp.integrate(psi[i]*f, (x, Omega[0], Omega[1]))
   c = A.LUsolve(b)
   u = 0
   for i in range(len(psi)):
       u += c[i,0]*psi[i]
   return u, c
```

Observe that we exploit the symmetry of the coefficient matrix: o upper triangular part is computed. Symbolic integration in sympy time consuming, and (roughly) halving the work has noticeable effect waiting time for the function to finish execution.

Fallback on numerical integration. Obviously, sympy mail fail to fully integrate $\int_{\Omega} \psi_i \psi_j dx$ and especially $\int_{\Omega} f \psi_i dx$ symbolically. There should extend the least_squares function such that it falls back on nu integration if the symbolic integration is unsuccessful. In the latter content returned value from sympy's integrate function is an object of type Ir We can test on this type and utilize the mpmath module in sympy to pumerical integration of high precision. Even when sympy manages to it symbolically, it can take an undesirable long time. We therefore incontent argument symbolic that governs whether or not to try symbolic integration of the improved version of function least_so

```
def least_squares(f, psi, Omega, symbolic=True):
    N = len(psi) - 1
    A = sp.zeros((N+1, N+1))
    b = sp.zeros((N+1, 1))
    x = sp.Svmbol('x')
    for i in range(N+1):
        for j in range(i, N+1):
            integrand = psi[i]*psi[j]
            if symbolic:
                I = sp.integrate(integrand, (x, Omega[0], Omega[1
            if not symbolic or isinstance(I, sp.Integral):
                # Could not integrate symbolically,
                # fall back on numerical integration
                integrand = sp.lambdify([x], integrand)
                I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]
            A[i,j] = \hat{A}[j,i] = \hat{I}
        integrand = psi[i]*f
        if symbolic:
            I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
        if not symbolic or isinstance(I, sp.Integral):
            integrand = sp.lambdify([x], integrand)
            I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]])
        b[i,0] = I
    c = A.LUsolve(b) # symbolic solve
    # c is a sympy Matrix object, numbers are in c[i,0]
    u = sum(c[i,0]*psi[i] for i in range(len(psi)))
    return u, [c[i,0] for i in range(len(c))]
```

The function is found in the file approx1D.py.

Plotting the approximation. Comparing the given f(x) and the imate u(x) visually is done by the following function, which with the sympy's lambdify tool converts a sympy expression to a Python func numerical computations:

```
lef comparison_plot(f, u, Omega, filename='tmp.pdf'):
    x = sp.Symbol('x')
    f = sp.lambdify([x], f, modules="numpy")
    u = sp.lambdify([x], u, modules="numpy")
    resolution = 401 # no of points in plot
    xcoor = linspace(Omega[0], Omega[1], resolution)
    exact = f(xcoor)
    approx = u(xcoor)
    plot(xcoor, approx)
    hold('on')
    plot(xcoor, exact)
    legend(['approximation', 'exact'])
    savefig(filename)
```

he modules='numpy' argument to lambdify is important if there are mathenatical functions, such as sin or exp in the symbolic expressions in f or u, and nese mathematical functions are to be used with vector arguments, like xcoor bove.

Both the least_squares and comparison_plot are found and coded in ne file approx1D.py². The forthcoming examples on their use appear in $x_approx1D.py$.

.5 Perfect approximation

et us use the code above to recompute the problem from Section 2.3 where we ant to approximate a parabola. What happens if we add an element x^2 to the asis and test what the best approximation is if V is the space of all parabolic inctions? The answer is quickly found by running

```
>>> from approx1D import *
>>> x = sp.Symbol('x')
>>> f = 10*(x-1)**2-1
>>> u, c = least_squares(f=f, psi=[1, x, x**2], Omega=[1, 2])
>>> print u
l0*x**2 - 20*x + 9
>>> print sp.expand(f)
l0*x**2 - 20*x + 9
```

Now, what if we use $\psi_i(x) = x^i$ for i = 0, 1, ..., N = 40? The output from east_squares gives $c_i = 0$ for i > 2, which means that the method finds the erfect approximation.

In fact, we have a general result that if $f \in V$, the least squares and rojection/Galerkin methods compute the exact solution u = f. The proof is raightforward: if $f \in V$, f can be expanded in terms of the basis functions, $= \sum_{j \in \mathcal{I}_s} d_j \psi_j$, for some coefficients $\{d_i\}_{i \in \mathcal{I}_s}$, and the right-hand side then has atries

$$b_i = (f, \psi_i) = \sum_{j \in \mathcal{I}_s} d_j(\psi_j, \psi_i) = \sum_{j \in \mathcal{I}_s} d_j A_{i,j}.$$

The linear system $\sum_{j} A_{i,j} c_j = b_i$, $i \in \mathcal{I}_s$, is then

$$\sum_{j \in \mathcal{I}_s} c_j A_{i,j} = \sum_{j \in \mathcal{I}_s} d_j A_{i,j}, \quad i \in \mathcal{I}_s,$$

which implies that $c_i = d_i$ for $i \in \mathcal{I}_s$.

2.6 Ill-conditioning

The computational example in Section 2.5 applies the least_squares is which invokes symbolic methods to calculate and solve the linear syste correct solution $c_0 = 9$, $c_1 = -20$, $c_2 = 10$, $c_i = 0$ for $i \ge 3$ is perfectly re

Suppose we convert the matrix and right-hand side to floating-poin and then solve the system using finite-precision arithmetics, which is w will (almost) always do in real life. This time we get astonishing resu to about N=7 we get a solution that is reasonably close to the ex Increasing N shows that seriously wrong coefficients are computed. If a table showing the solution of the linear system arising from approx a parabola by functions on the form $u(x) = c_0 + c_1 x + c_2 x^2 + \cdots +$ Analytically, we know that $c_j = 0$ for j > 2, but numerically we may ge for j > 2.

exact	sympy	numpy32	numpy64
9	9.62	5.57	8.98
-20	-23.39	-7.65	-19.93
10	17.74	-4.50	9.96
0	-9.19	4.13	-0.26
0	5.25	2.99	0.72
0	0.18	-1.21	-0.93
0	-2.48	-0.41	0.73
0	1.81	-0.013	-0.36
0	-0.66	0.08	0.11
0	0.12	0.04	-0.02
0	-0.001	-0.02	0.002

The exact value of c_j , j = 0, 1, ..., 10, appears in the first column w other columns correspond to results obtained by three different method

- Column 2: The matrix and vector are converted to the data st sympy.mpmath.fp.matrix and the sympy.mpmath.fp.lu_solve is used to solve the system.
- Column 3: The matrix and vector are converted to numpy arrays w type numpy.float32 (single precision floating-point number) and by the numpy.linalg.solve function.
- Column 4: As column 3, but the data type is numpy.float64 precision floating-point number).

²http://tinyurl.com/nm5587k/fem/approx1D.py

/e see from the numbers in the table that double precision performs much better nan single precision. Nevertheless, when plotting all these solutions the curves annot be visually distinguished (!). This means that the approximations look erfect, despite the partially very wrong values of the coefficients.

Increasing N to 12 makes the numerical solver in numpy abort with the lessage: "matrix is numerically singular". A matrix has to be non-singular to be invertible, which is a requirement when solving a linear system. Already when he matrix is close to singular, it is ill-conditioned, which here implies that the umerical solution algorithms are sensitive to round-off errors and may produce very) inaccurate results.

The reason why the coefficient matrix is nearly singular and ill-conditioned that our basis functions $\psi_i(x)=x^i$ are nearly linearly dependent for large i. hat is, x^i and x^{i+1} are very close for i not very small. This phenomenon is lustrated in Figure 4. There are 15 lines in this figure, but only half of them re visually distinguishable. Almost linearly dependent basis functions give rise an ill-conditioned and almost singular matrix. This fact can be illustrated by imputing the determinant, which is indeed very close to zero (recall that a zero eterminant implies a singular and non-invertible matrix): 10^{-65} for N=10 and 10^{-92} for 10^{-92} for

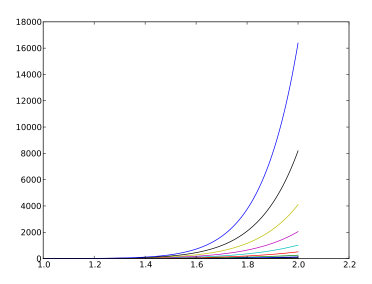


Figure 4: The 15 first basis functions x^i , i = 0, ..., 14.

On the other hand, the double precision numpy solver do run for N = 100, sulting in answers that are not significantly worse than those in the table

above, and large powers are associated with small coefficients (e.g., c_j for $10 \le j \le 20$ and $c < 10^{-5}$ for j > 20). Even for N = 100 the approx still lies on top of the exact curve in a plot (!).

The conclusion is that visual inspection of the quality of the approx may not uncover fundamental numerical problems with the computation ever, numerical analysts have studied approximations and ill-condition decades, and it is well known that the basis $\{1, x, x^2, x^3, \ldots, \}$ is a be The best basis from a matrix conditioning point of view is to have ort functions such that $(\psi_i, \psi_j) = 0$ for $i \neq j$. There are many known sets of onal polynomials and other functions. The functions used in the finite methods are almost orthogonal, and this property helps to avoid problems olving matrix systems. Almost orthogonal is helpful, but not enoug it comes to partial differential equations, and ill-conditioning of the comparison of the comparison of the comparison of the solving large-scale matrix systems arising from element discretizations.

2.7 Fourier series

A set of sine functions is widely used for approximating functions (the s also orthogonal as explained more in Section 2.6). Let us take

$$V = \operatorname{span} \left\{ \sin \pi x, \sin 2\pi x, \dots, \sin (N+1)\pi x \right\}.$$

That is,

$$\psi_i(x) = \sin((i+1)\pi x), \quad i \in \mathcal{I}_s.$$

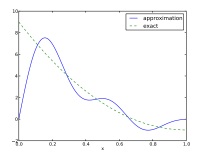
An approximation to the f(x) function from Section 2.3 can then be co by the least_squares function from Section 2.4:

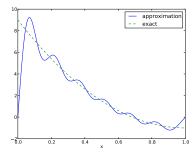
```
N = 3
from sympy import sin, pi
x = sp.Symbol('x')
psi = [sin(pi*(i+1)*x) for i in range(N+1)]
f = 10*(x-1)**2 - 1
Omega = [0, 1]
u, c = least_squares(f, psi, Omega)
comparison_plot(f, u, Omega)
```

Figure 5 (left) shows the oscillatory approximation of $\sum_{j=0}^{N} c_j \sin((j \text{ when } N = 3$. Changing N to 11 improves the approximation considers Figure 5 (right).

There is an error f(0) - u(0) = 9 at x = 0 in Figure 5 regardless of he N is, because all $\psi_i(0) = 0$ and hence u(0) = 0. We may help the approx to be correct at x = 0 by seeking

$$u(x) = f(0) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x) .$$





igure 5: Best approximation of a parabola by a sum of 3 (left) and 11 (right) ne functions.

lowever, this adjustment introduces a new problem at x = 1 since we now get a error f(1) - u(1) = f(1) - 0 = -1 at this point. A more clever adjustment is a replace the f(0) term by a term that is f(0) at x = 0 and f(1) at x = 1. A mple linear combination f(0)(1-x) + xf(1) does the job:

$$u(x) = f(0)(1-x) + xf(1) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x).$$
 (40)

his adjustment of u alters the linear system slightly. In the general case, we set

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x),$$

nd the linear system becomes

$$\sum_{j \in \mathcal{I}_s} (\psi_i, \psi_j) c_j = (f - B, \psi_i), \quad i \in \mathcal{I}_s.$$

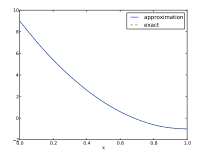
he calculations can still utilize the least_squares or least_squares_orth unctions, but solve for u-b:

```
f0 = 0; f1 = -1
3 = f0*(1-x) + x*f1
1_sum, c = least_squares_orth(f-b, psi, Omega)
1 = B + u_sum
```

Figure 6 shows the result of the technique for ensuring right boundary values. ven 3 sines can now adjust the f(0)(1-x)+xf(1) term such that u approximates 12 parabola really well, at least visually.

.8 Orthogonal basis functions

he choice of sine functions $\psi_i(x) = \sin((i+1)\pi x)$ has a great computational dvantage: on $\Omega = [0,1]$ these basis functions are *orthogonal*, implying that i,j=0 if $i \neq j$. This result is realized by trying



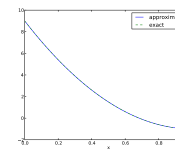


Figure 6: Best approximation of a parabola by a sum of 3 (left) and 1: sine functions with a boundary term.

```
integrate(sin(j*pi*x)*sin(k*pi*x), x, 0, 1)
```

in WolframAlpha³ (avoid i in the integrand as this symbol means the in unit $\sqrt{-1}$). Also by asking WolframAlpha about $\int_0^1 \sin^2(j\pi x) dx$, we fi equal 1/2. With a diagonal matrix we can easily solve for the coefficient hand:

$$c_i = 2 \int_0^1 f(x) \sin((i+1)\pi x) dx, \quad i \in \mathcal{I}_s,$$

which is nothing but the classical formula for the coefficients of the Fourier series of f(x) on [0,1]. In fact, when V contains the basic functions us Fourier series expansion, the approximation method derived in Section 2 in the classical Fourier series for f(x) (see Exercise 8 for details).

With orthogonal basis functions we can make the $least_squares$ f (much) more efficient since we know that the matrix is diagonal and ϵ diagonal elements need to be computed:

```
def least_squares_orth(f, psi, Omega):
    N = len(psi) - 1
    A = [0]*(N+1)
    b = [0]*(N+1)
    x = sp.Symbol('x')
    for i in range(N+1):
        A[i] = sp.integrate(psi[i]**2, (x, Omega[0], Omega[1]))
        b[i] = sp.integrate(psi[i]*f, (x, Omega[0], Omega[1]))
    c = [b[i]/A[i] for i in range(len(b))]
    u = 0
    for i in range(len(psi)):
        u += c[i]*psi[i]
    return u, c
```

As mentioned in Section 2.4, symbolic integration may fail or take vertime. It is therefore natural to extend the implementation above with a

³http://wolframalpha.com

here we can choose between symbolic and numerical integration and fall back a the latter if the former fails:

```
lef least_squares_orth(f, psi, Omega, symbolic=True):
   N = len(psi) - 1
   A = \lceil 0 \rceil * (N+1)
                       # plain list to hold symbolic expressions
   b = [0]*(N+1)
   x = sp.Symbol('x')
   for i in range(N+1):
       # Diagonal matrix term
       A[i] = sp.integrate(psi[i]**2, (x, Omega[0], Omega[1]))
       # Right-hand side term
       integrand = psi[i]*f
       if symbolic:
           I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
       if not symbolic or isinstance(I, sp.Integral):
           print 'numerical integration of', integrand
           integrand = sp.lambdify([x], integrand)
           I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]])
       b[i] = I
   c = [b[i]/A[i] for i in range(len(b))]
   u = 0
   u = sum(c[i,0]*psi[i] for i in range(len(psi)))
   return u. c
```

his function is found in the file approx1D.py. Observe that we here assume at $\int_{\Omega} \varphi_i^2 dx$ can always be symbolically computed, which is not an unreasonable sumption when the basis functions are orthogonal, but there is no guarantee, so a improved version of the function above would implement numerical integration lso for the A[i,i] term.

.9 Numerical computations

ometimes the basis functions ψ_i and/or the function f have a nature that takes symbolic integration CPU-time consuming or impossible. Even though we nplemented a fallback on numerical integration of $\int f \varphi_i dx$ considerable time light be required by sympy in the attempt to integrate symbolically. Therefore, will be handy to have function for fast numerical integration and numerical plution of the linear system. Below is such a method. It requires Python unctions f(x) and psi(x,i) for f(x) and $\psi_i(x)$ as input. The output is a mesh unction with values u on the mesh with points in the array x. Three numerical itegration methods are offered: scipy.integrate.quad (precision set to 10^{-8}), ympy.mpmath.quad (high precision), and a Trapezoidal rule based on the points x.

```
dx = x[1] - x[0]
   for i in range(N+1):
        j limit = i+1 if orthogonal basis else N+1
        for j in range(i, j_limit):
            print '(%d,%d)' % (i, j)
            if integration method == 'scipy':
                A_ij = scipy.integrate.quad(
                    lambda x: psi(x,i)*psi(x,j),
                    Omega[0], Omega[1], epsabs=1E-9, epsrel=1E-9)
            elif integration method == 'sympy':
                A_ij = sp.mpmath.quad(
                    lambda x: psi(x,i)*psi(x,j),
                    [Omega[0], Omega[1]])
            else:
                values = psi(x,i)*psi(x,j)
                A_ij = trapezoidal(values, dx)
            A[i,j] = A[j,i] = A_{ij}
        if integration_method == 'scipy':
            b_i = scipy.integrate.quad(
                lambda x: f(x)*psi(x,i), Omega[0], Omega[1],
                epsabs=1E-9, epsrel=1E-9)[0]
        elif integration_method == 'sympy':
            b_i = sp.mpmath.quad(
                lambda x: f(x)*psi(x,i), [Omega[0], Omega[1]])
            values = f(x)*psi(x,i)
            b i = trapezoidal(values, dx)
        b[i] = b i
    c = b/np.diag(A) if orthogonal_basis else np.linalg.solve(A,
    u = sum(c[i]*psi(x, i) for i in range(N+1))
   return u, c
def trapezoidal(values, dx):
    """Integrate values by the Trapezoidal rule (mesh size dx).""
    return dx*(np.sum(values) - 0.5*values[0] - 0.5*values[-1])
```

Here is an example on calling the function:

2.10 The interpolation (or collocation) method

The principle of minimizing the distance between u and f is an intuit of computing a best approximation $u \in V$ to f. However, there as approaches as well. One is to demand that $u(x_i) = f(x_i)$ at some selecte x_i , $i \in \mathcal{I}_s$:

$$u(x_i) = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x_i) = f(x_i), \quad i \in \mathcal{I}_s.$$
 (42)

his criterion also gives a linear system with N+1 unknown coefficients $\{c_i\}_{i\in\mathcal{I}}$:

$$\sum_{j \in \mathcal{I}_s} A_{i,j} c_j = b_i, \quad i \in \mathcal{I}_s, \tag{43}$$

ith

$$A_{i,j} = \psi_j(x_i), \tag{44}$$

$$b_i = f(x_i). (45)$$

his time the coefficient matrix is not symmetric because $\psi_j(x_i) \neq \psi_i(x_j)$ in eneral. The method is often referred to as an interpolation method since some oint values of f are given $(f(x_i))$ and we fit a continuous function u that goes rough the $f(x_i)$ points. In this case the x_i points are called interpolation vints. When the same approach is used to approximate differential equations, ne usually applies the name collocation method and x_i are known as collocation vints.

Given f as a sympy symbolic expression f, $\{\psi_i\}_{i\in\mathcal{I}_s}$ as a list psi, and a set f points $\{x_i\}_{i\in\mathcal{I}_s}$ as a list or array points, the following Python function sets f and solves the matrix system for the coefficients $\{c_i\}_{i\in\mathcal{I}_s}$:

```
lef interpolation(f, psi, points):
   N = len(psi) - 1
   A = sp.zeros((N+1, N+1))
   b = sp.zeros((N+1, 1))
   x = sp.Symbol('x')
   # Turn psi and f into Python functions
   psi = [sp.lambdify([x], psi[i]) for i in range(N+1)]
   f = sp.lambdify([x], f)
   for i in range(N+1):
       for j in range(N+1):
           A[i,j] = psi[j](points[i])
       b[i,0] = f(points[i])
   c = A.LUsolve(b)
   11 = 0
   for i in range(len(psi)):
       u += c[i,0]*psi[i](x)
   return u
```

he interpolation function is a part of the ${\tt approx1D}$ module.

We found it convenient in the above function to turn the expressions f nd psi into ordinary Python functions of x, which can be called with float alues in the list points when building the matrix and the right-hand side. he alternative is to use the subs method to substitute the x variable in an approaches in a simple setting:

```
>>> from sympy import *
>>> x = Symbol('x')
>>> e = x**2
                          # symbolic expression involving x
>>> p = 0.5
                          # a value of x
>>> v = e.subs(x, p)
                          # evaluate e for x=p
0.250000000000000
>>> type(v)
sympy.core.numbers.Float
>>> e = lambdify([x], e) # make Python function of e
>>> type(e)
>>> function
                          # evaluate e(x) for x=p
>>> v = e(p)
>>> v
0.25
>>> type(v)
float
```

A nice feature of the interpolation or collocation method is that it computing integrals. However, one has to decide on the location of the x A simple, yet common choice, is to distribute them uniformly throughout

Example. Let us illustrate the interpolation or collocation method by imating our parabola $f(x) = 10(x-1)^2 - 1$ by a linear function on Ω using two collocation points $x_0 = 1 + 1/3$ and $x_1 = 1 + 2/3$:

```
f = 10*(x-1)**2 - 1
psi = [1, x]
Omega = [1, 2]
points = [1 + sp.Rational(1,3), 1 + sp.Rational(2,3)]
u = interpolation(f, psi, points)
comparison_plot(f, u, Omega)
```

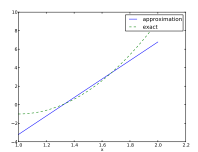
The resulting linear system becomes

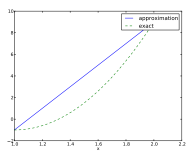
$$\left(\begin{array}{cc} 1 & 4/3 \\ 1 & 5/3 \end{array}\right) \left(\begin{array}{c} c_0 \\ c_1 \end{array}\right) = \left(\begin{array}{c} 1/9 \\ 31/9 \end{array}\right)$$

with solution $c_0 = -119/9$ and $c_1 = 10$. Figure 7 (left) shows the rapproximation u = -119/9 + 10x. We can easily test other interpolation say $x_0 = 1$ and $x_1 = 2$. This changes the line quite significantly, see I (right).

2.11 Lagrange polynomials

In Section 2.7 we explain the advantage with having a diagonal matrix: f for the coefficients $\{c_i\}_{i\in\mathcal{I}_s}$ can then be derived by hand. For an interpolation method a diagonal matrix implies that $\psi_j(x_i) = 0$ if $i \neq 1$ set of basis functions $\psi_i(x)$ with this property is the Lagrange interpolynomials, or just Lagrange polynomials. (Although the functions are after Lagrange, they were first discovered by Waring in 1779, rediscovered by Waring in 1779, rediscovered





igure 7: Approximation of a parabola by linear functions computed by two terpolation points: 4/3 and 5/3 (left) versus 1 and 2 (right).

uler in 1783, and published by Lagrange in 1795.) The Lagrange polynomials ave the form

$$\psi_i(x) = \prod_{j=0, j \neq i}^{N} \frac{x - x_j}{x_i - x_j} = \frac{x - x_0}{x_i - x_0} \cdots \frac{x - x_{i-1}}{x_i - x_{i-1}} \frac{x - x_{i+1}}{x_i - x_{i+1}} \cdots \frac{x - x_N}{x_i - x_N}, \quad (46)$$

or $i \in \mathcal{I}_s$. We see from (46) that all the ψ_i functions are polynomials of degree ℓ which have the property

$$\psi_i(x_s) = \delta_{is}, \quad \delta_{is} = \begin{cases} 1, & i = s, \\ 0, & i \neq s, \end{cases}$$

$$\tag{47}$$

hen x_s is an interpolation/collocation point. Here we have used the *Kronecker elta* symbol δ_{is} . This property implies that $A_{i,j} = 0$ for $i \neq j$ and $A_{i,j} = 1$ hen i = j. The solution of the linear system is them simply

$$c_i = f(x_i), \quad i \in \mathcal{I}_s,$$
 (48)

nd

$$u(x) = \sum_{j \in \mathcal{I}_s} f(x_i)\psi_i(x). \tag{49}$$

The following function computes the Lagrange interpolating polynomial $\psi_i(x)$, iven the interpolation points x_0, \ldots, x_N in the list or array points:

```
lef Lagrange_polynomial(x, i, points):
    p = 1
    for k in range(len(points)):
        if k != i:
        p *= (x - points[k])/(points[i] - points[k])
    return p
```

he next function computes a complete basis using equidistant points throughout $\dot{}$

```
def Lagrange_polynomials_01(x, N):
    if isinstance(x, sp.Symbol):
        h = sp.Rational(1, N-1)
    else:
        h = 1.0/(N-1)
    points = [i*h for i in range(N)]
    psi = [Lagrange_polynomial(x, i, points) for i in range(N)]
    return psi, points
```

When x is an sp.Symbol object, we let the spacing between the interproints, h, be a sympy rational number for nice end results in the for ψ_i . The other case, when x is a plain Python float, signifies nu computing, and then we let h be a floating-point number. Observe t Lagrange_polynomial function works equally well in the symbolic and m case - just think of x being an sp.Symbol object or a Python float. interactive session illustrates the difference between symbolic and nu computing of the basis functions and points:

```
>>> import sympy as sp
>>> x = sp.Symbol('x')
>>> psi, points = Lagrange_polynomials_01(x, N=3)
>>> points
[0, 1/2, 1]
>>> psi
[(1 - x)*(1 - 2*x), 2*x*(2 - 2*x), -x*(1 - 2*x)]
>>> x = 0.5  # numerical computing
>>> psi, points = Lagrange_polynomials_01(x, N=3)
>>> points
[0.0, 0.5, 1.0]
>>> psi
[-0.0, 1.0, 0.0]
```

The Lagrange polynomials are very much used in finite element methods of their property (47).

Approximation of a polynomial. The Galerkin or least squares metl to an exact approximation if f lies in the space spanned by the basis function be interest to see how the interpolation method with Lagrange poly as basis is able to approximate a polynomial, e.g., a parabola. Running

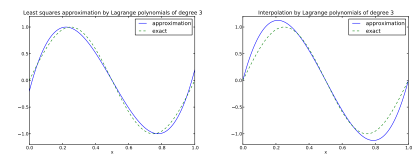
```
for N in 2, 4, 5, 6, 8, 10, 12:
    f = x**2
    psi, points = Lagrange_polynomials_01(x, N)
    u = interpolation(f, psi, points)
```

shows the result that up to N=4 we achieve an exact approximation, a round-off errors start to grow, such that N=15 leads to a 15-degree poly for u where the coefficients in front of x^r for r > 2 are of size 10^{-5} and

Successful example. Trying out the Lagrange polynomial basis for mating $f(x) = \sin 2\pi x$ on $\Omega = [0, 1]$ with the least squares and the interjetechniques can be done by

```
c = sp.Symbol('x')
f = sp.sin(2*sp.pi*x)
psi, points = Lagrange_polynomials_01(x, N)
lmega=[0, 1]
1, c = least_squares(f, psi, Omega)
comparison_plot(f, u, Omega)
1, c = interpolation(f, psi, points)
comparison_plot(f, u, Omega)
```

igure 8 shows the results. There is little difference between the least squares and n interpolation technique. Increasing N gives visually better approximations.



igure 8: Approximation via least squares (left) and interpolation (right) of a ne function by Lagrange interpolating polynomials of degree 3.

ess successful example. The next example concerns interpolating f(x) = -2x| on $\Omega = [0,1]$ using Lagrange polynomials. Figure 9 shows a peculiar fect: the approximation starts to oscillate more and more as N grows. This umerical artifact is not surprising when looking at the individual Lagrange olynomials. Figure 10 shows two such polynomials, $\psi_2(x)$ and $\psi_7(x)$, both of egree 11 and computed from uniformly spaced points $x_{x_i} = i/11$, $i = 0, \ldots, 11$, tarked with circles. We clearly see the property of Lagrange polynomials: $2(x_i) = 0$ and $\psi_7(x_i) = 0$ for all i, except $\psi_2(x_2) = 1$ and $\psi_7(x_7) = 1$. The lost striking feature, however, is the significant oscillation near the boundary, he reason is easy to understand: since we force the functions to zero at so many oints, a polynomial of high degree is forced to oscillate between the points. The henomenon is named Runge's phenomenon and you can read a more detailed splanation on Wikipedia⁴.

temedy for strong oscillations. The oscillations can be reduced by a more ever choice of interpolation points, called the *Chebyshev nodes*:

$$x_i = \frac{1}{2}(a+b) + \frac{1}{2}(b-a)\cos\left(\frac{2i+1}{2(N+1)}pi\right), \quad i = 0..., N,$$
 (50)

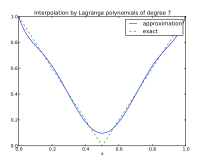
on the interval $\Omega = [a,b]$. Here is a flexible version of the Lagrange_polyn function above, valid for any interval $\Omega = [a,b]$ and with the possibility t ate both uniformly distributed points and Chebyshev nodes:

```
def Lagrange_polynomials(x, N, Omega, point_distribution='uniform
   if point_distribution == 'uniform':
        if isinstance(x, sp.Symbol):
            h = sp.Rational(Omega[1] - Omega[0], N)
        else:
            h = (Omega[1] - Omega[0])/float(N)
        points = [Omega[0] + i*h for i in range(N+1)]
   elif point_distribution == 'Chebyshev':
        points = Chebyshev_nodes(Omega[0], Omega[1], N)
   psi = [Lagrange_polynomial(x, i, points) for i in range(N+1)]
   return psi, points

def Chebyshev_nodes(a, b, N):
   from math import cos, pi
   return [0.5*(a+b) + 0.5*(b-a)*cos(float(2*i+1)/(2*N+1))*pi) \
        for i in range(N+1)]
```

All the functions computing Lagrange polynomials listed above are foun module file Lagrange.py. Figure 11 shows the improvement of using Ch nodes (compared with Figure 9). The reason is that the corresponding L polynomials have much smaller oscillations as seen in Figure 12 (compa Figure 10).

Another cure for undesired oscillation of higher-degree interpolating nomials is to use lower-degree Lagrange polynomials on many small patter domain, which is the idea pursued in the finite element method, stance, linear Lagrange polynomials on [0,1/2] and [1/2,1] would yield a approximation to f(x) = |1 - 2x| on $\Omega = [0,1]$ since f is piecewise linear



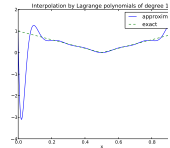
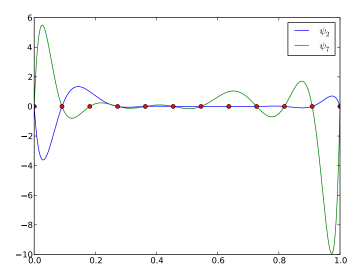


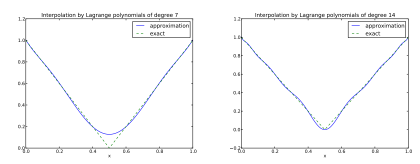
Figure 9: Interpolation of an absolute value function by Lagrange poly and uniformly distributed interpolation points: degree 7 (left) and 14 (

How does the least squares or projection methods work with Lapolynomials? We can just call the least_squares function, but symproblems integrating the f(x) = |1 - 2x| function times a polynomianeed to fallback on numerical integration.

⁴http://en.wikipedia.org/wiki/Runge%27s phenomenon



igure 10: Illustration of the oscillatory behavior of two Lagrange polynomials ased on 12 uniformly spaced points (marked by circles).



igure 11: Interpolation of an absolute value function by Lagrange polynomials nd Chebyshev nodes as interpolation points: degree 7 (left) and 14 (right).

```
lef least_squares(f, psi, Omega):
    N = len(psi) - 1
    A = sp.zeros((N+1, N+1))
    b = sp.zeros((N+1, 1))
    x = sp.Symbol('x')
    for i in range(N+1):
        for j in range(i, N+1):
            integrand = psi[i]*psi[j]
            I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
            if isinstance(I, sp.Integral):
```

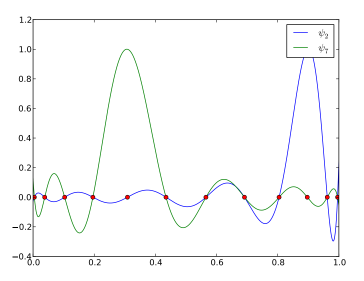


Figure 12: Illustration of the less oscillatory behavior of two Lagrange mials based on 12 Chebyshev points (marked by circles).

```
# Could not integrate symbolically, fallback
            # on numerical integration with mpmath.quad
            integrand = sp.lambdify([x], integrand)
            I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]
        A[i,j] = \overline{A}[j,i] = \overline{I}
    integrand = psi[i]*f
    I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
    if isinstance(I, sp.Integral):
        integrand = sp.lambdify([x], integrand)
        I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]])
    b[i,0] = I
c = A.LUsolve(b)
u = 0
for i in range(len(psi)):
    u += c[i,0]*psi[i]
return u
```

3 Finite element basis functions

The specific basis functions exemplified in Section 2 are in general nonzer entire domain Ω , see Figure 13 for an example where we plot $\psi_0(x) =$ and $\psi_1(x) = \sin 2\pi x$ together with a possible sum $u(x) = 4\psi_0(x) -$ We shall now turn the attention to basis functions that have *compact* meaning that they are nonzero on only a small portion of Ω . Moreo

nall restrict the functions to be *piecewise polynomials*. This means that the omain is split into subdomains and the function is a polynomial on one or more abdomains, see Figure 14 for a sketch involving locally defined hat functions nat make $u = \sum_j c_j \psi_j$ piecewise linear. At the boundaries between subdomains ne normally forces continuity of the function only so that when connecting two olynomials from two subdomains, the derivative becomes discontinuous. These the of basis functions are fundamental in the finite element method.

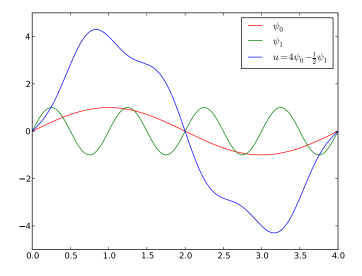


Figure 13: A function resulting from adding two sine basis functions.

We first introduce the concepts of elements and nodes in a simplistic fashion s often met in the literature. Later, we shall generalize the concept of an ement, which is a necessary step to treat a wider class of approximations within 12 family of finite element methods. The generalization is also compatible with 13 necessary used in the FEniCS⁵ finite element software.

.1 Elements and nodes

et us divide the interval Ω on which f and u are defined into non-overlapping ibintervals $\Omega^{(e)}$, $e = 0, \dots, N_e$:

$$\Omega = \Omega^{(0)} \cup \dots \cup \Omega^{(N_e)} \,. \tag{51}$$

We shall for now refer to $\Omega^{(e)}$ as an *element*, having number e. On each element e introduce a set of points called *nodes*. For now we assume that the nodes are

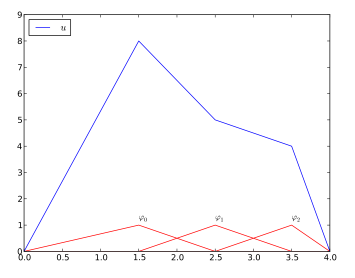


Figure 14: A function resulting from adding three local piecewise line functions.

uniformly spaced throughout the element and that the boundary point elements are also nodes. The nodes are given numbers both within an and in the global domain. These are referred to as *local* and *global* node respectively. Figure 15 shows element boundaries with small vertical line as small disks, element numbers in circles, and global node numbers ur nodes.

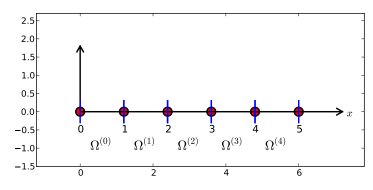


Figure 15: Finite element mesh with 5 elements and 6 nodes.

 $^{^5 {\}tt http://fenicsproject.org}$

Nodes and elements uniquely define a *finite element mesh*, which is our iscrete representation of the domain in the computations. A common special ase is that of a *uniformly partitioned mesh* where each element has the same eight and the distance between nodes is constant.

Example. On $\Omega = [0, 1]$ we may introduce two elements, $\Omega^{(0)} = [0, 0.4]$ and $S^{(1)} = [0.4, 1]$. Furthermore, let us introduce three nodes per element, equally paced within each element. Figure 16 shows the mesh. The three nodes in ement number 0 are $x_0 = 0$, $x_1 = 0.2$, and $x_2 = 0.4$. The local and global node umbers are here equal. In element number 1, we have the local nodes $x_0 = 0.4$, $x_1 = 0.7$, and $x_2 = 1$ and the corresponding global nodes $x_2 = 0.4$, $x_3 = 0.7$, and $x_4 = 1$. Note that the global node $x_2 = 0.4$ is shared by the two elements.

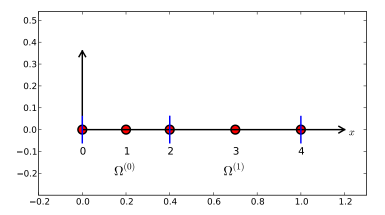


Figure 16: Finite element mesh with 2 elements and 5 nodes.

For the purpose of implementation, we introduce two lists or arrays: nodes or storing the coordinates of the nodes, with the global node numbers as indices, and elements for holding the global node numbers in each element, with the scal node numbers as indices. The nodes and elements lists for the sample tesh above take the form

```
nodes = [0, 0.2, 0.4, 0.7, 1]
elements = [[0, 1, 2], [2, 3, 4]]
```

ooking up the coordinate of local node number 2 in element 1 is here done by odes[elements[1][2]] (recall that nodes and elements start their numbering t 0).

The numbering of elements and nodes does not need to be regular. Figure 17 nows and example corresponding to

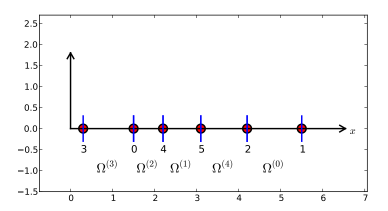


Figure 17: Example on irregular numbering of elements and node

3.2 The basis functions

Construction principles. Finite element basis functions are in this ognized by the notation $\varphi_i(x)$, where the index now in the beginning corr to a global node number. In the current approximation problem we shal take $\psi_i = \varphi_i$.

Let i be the global node number corresponding to local node r in number e. The finite element basis functions φ_i are now defined as follows:

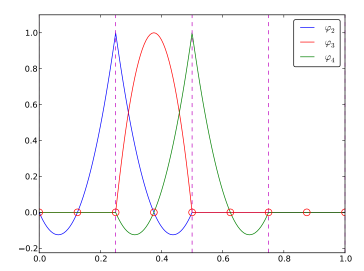
- If local node number r is not on the boundary of the element, tal to be the Lagrange polynomial that is 1 at the local node number zero at all other nodes in the element. On all other elements, φ_i :
- If local node number r is on the boundary of the element, let φ_i l up of the Lagrange polynomial over element e that is 1 at node i, or with the Lagrange polynomial over element e+1 that is also 1 at On all other elements, $\varphi_i = 0$.

A visual impression of three such basis functions are given in Figure 18

Properties of φ_i . The construction of basis functions according to th ples above lead to two important properties of $\varphi_i(x)$. First,

$$\varphi_i(x_j) = \delta_{ij}, \quad \delta_{ij} = \begin{cases} 1, & i = j, \\ 0, & i \neq j, \end{cases}$$

when x_j is a node in the mesh with global node number j. The result $\varphi_i(x)$ arises because the Lagrange polynomials are constructed to have exact property. The property also implies a convenient interpretation of c_i as to of u at node i. To show this, we expand u in the usual way as $\sum_j c_j$ choose $\psi_i = \varphi_i$:



igure 18: Illustration of the piecewise quadratic basis functions associated ith nodes in element 1.

$$u(x_i) = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x_i) = \sum_{j \in \mathcal{I}_s} c_j \varphi_j(x_i) = c_i \varphi_i(x_i) = c_i.$$

ecause of this interpretation, the coefficient c_i is by many named u_i or U_i . Second, $\varphi_i(x)$ is mostly zero throughout the domain:

- $\varphi_i(x) \neq 0$ only on those elements that contain global node i,
- $\varphi_i(x)\varphi_j(x)\neq 0$ if and only if i and j are global node numbers in the same element.

ince $A_{i,j}$ is the integral of $\varphi_i\varphi_j$ it means that most of the elements in the refficient matrix will be zero. We will come back to these properties and use nem actively in computations to save memory and CPU time.

We let each element have d+1 nodes, resulting in local Lagrange polynomials f degree d. It is not a requirement to have the same d value in each element, ut for now we will assume so.

.3 Example on piecewise quadratic finite element functions

igure 18 illustrates how piecewise quadratic basis functions can look like (d=2). /e work with the domain $\Omega = [0,1]$ divided into four equal-sized elements, each

having three nodes. The nodes and elements lists in this particular ϵ become

```
nodes = [0, 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875, 1.0]
elements = [[0, 1, 2], [2, 3, 4], [4, 5, 6], [6, 7, 8]]
```

Figure 19 sketches the mesh and the numbering. Nodes are marked with on the x axis and element boundaries are marked with vertical dashed Figure 18.

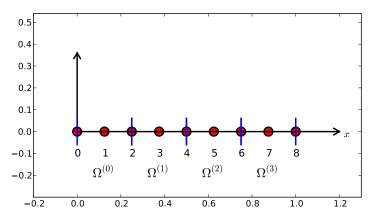


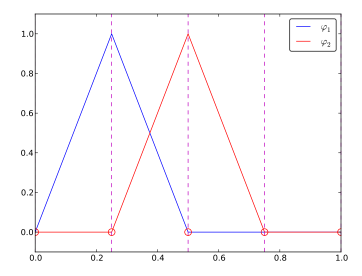
Figure 19: Sketch of mesh with 4 elements and 3 nodes per eleme

Let us explain in detail how the basis functions are constructed act to the principles. Consider element number 1 in Figure 18, $\Omega^{(1)} = [0]$ with local nodes 0, 1, and 2 corresponding to global nodes 2, 3, and coordinates of these nodes are 0.25, 0.375, and 0.5, respectively. We define Lagrange polynomials on this element:

- 1. The polynomial that is 1 at local node 1 (x = 0.375, global node 3 up the basis function $\varphi_3(x)$ over this element, with $\varphi_3(x) = 0$ out element.
- 2. The Lagrange polynomial that is 1 at local node 0 is the "rigl of the global basis function $\varphi_2(x)$. The "left part" of $\varphi_2(x)$ cor a Lagrange polynomial associated with local node 2 in the neig element $\Omega^{(0)} = [0, 0.25]$.
- 3. Finally, the polynomial that is 1 at local node 2 (global node 4) is part" of the global basis function $\varphi_4(x)$. The "right part" comes f Lagrange polynomial that is 1 at local node 0 in the neighboring $\Omega^{(2)} = [0.5, 0.75]$.

As mentioned earlier, any global basis function $\varphi_i(x)$ is zero on element do not contain the node with global node number i.

The other global functions associated with internal nodes, φ_1 , φ_5 , and φ_7 , are ll of the same shape as the drawn φ_3 , while the global basis functions associated ith shared nodes also have the same shape, provided the elements are of the ame length.



igure 20: Illustration of the piecewise linear basis functions associated with odes in element 1.

.4 Example on piecewise linear finite element functions

igure 20 shows piecewise linear basis functions (d=1). Also here we have sur elements on $\Omega=[0,1]$. Consider the element $\Omega^{(1)}=[0.25,0.5]$. Now there re no internal nodes in the elements so that all basis functions are associated ith nodes at the element boundaries and hence made up of two Lagrange olynomials from neighboring elements. For example, $\varphi_1(x)$ results from the agrange polynomial in element 0 that is 1 at local node 1 and 0 at local node , combined with the Lagrange polynomial in element 1 that is 1 at local node 0 at local node 1. The other basis functions are constructed similarly.

Explicit mathematical formulas are needed for $\varphi_i(x)$ in computations. In the iecewise linear case, one can show that

$$\varphi_{i}(x) = \begin{cases} 0, & x < x_{i-1}, \\ (x - x_{i-1})/(x_{i} - x_{i-1}), & x_{i-1} \le x < x_{i}, \\ 1 - (x - x_{i})/(x_{i+1} - x_{i}), & x_{i} \le x < x_{i+1}, \\ 0, & x \ge x_{i+1}. \end{cases}$$
(53)

Here, x_j , j = i - 1, i, i + 1, denotes the coordinate of node j. For elen equal length h the formulas can be simplified to

$$\varphi_i(x) = \begin{cases} 0, & x < x_{i-1}, \\ (x - x_{i-1})/h, & x_{i-1} \le x < x_i, \\ 1 - (x - x_i)/h, & x_i \le x < x_{i+1}, \\ 0, & x \ge x_{i+1} \end{cases}$$

3.5 Example on piecewise cubic finite element basis tions

Piecewise cubic basis functions can be defined by introducing four no element. Figure 21 shows examples on $\varphi_i(x)$, i = 3, 4, 5, 6, associate element number 1. Note that φ_4 and φ_5 are nonzero on element number φ_3 and φ_6 are made up of Lagrange polynomials on two neighboring elements.

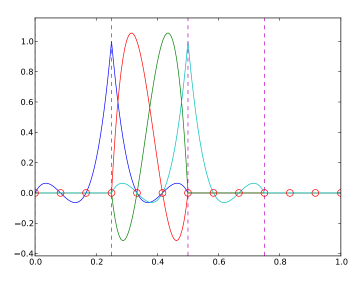


Figure 21: Illustration of the piecewise cubic basis functions associat nodes in element 1.

We see that all the piecewise linear basis functions have the same "hat They are naturally referred to as hat functions, also called chapeau fu. The piecewise quadratic functions in Figure 18 are seen to be of tw "Rounded hats" associated with internal nodes in the elements and son "sombrero" shaped hats associated with element boundary nodes. High basis functions also have hat-like shapes, but the functions have proroscillations in addition, as illustrated in Figure 21.

A common terminology is to speak about *linear elements* as elements with two scal nodes associated with piecewise linear basis functions. Similarly, *quadratic lements* and *cubic elements* refer to piecewise quadratic or cubic functions ver elements with three or four local nodes, respectively. Alternative names, equently used later, are P1 elements for linear elements, P2 for quadratic ements, and so forth: Pd signifies degree d of the polynomial basis functions.

.6 Calculating the linear system

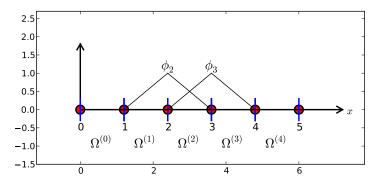
he elements in the coefficient matrix and right-hand side are given by the rmulas (27) and (28), but now the choice of ψ_i is φ_i . Consider P1 elements here $\varphi_i(x)$ piecewise linear. Nodes and elements numbered consecutively from ft to right in a uniformly partitioned mesh imply the nodes

$$x_i = ih, \quad i = 0, \dots, N,$$

nd the elements

$$\Omega^{(i)} = [x_i, x_{i+1}] = [ih, (i+1)h], \quad i = 0, \dots, N_e = N - 1.$$
 (55)

We have in this case N elements and N+1 nodes, and $\Omega=[x_0,x_N]$. The rmula for $\varphi_i(x)$ is given by (54) and a graphical illustration is provided in igures 20 and 23. First we clearly see from the figures the very important roperty $\varphi_i(x)\varphi_j(x)\neq 0$ if and only if $j=i-1,\ j=i,$ or j=i+1, or lternatively expressed, if and only if i and j are nodes in the same element. Therwise, φ_i and φ_j are too distant to have an overlap and consequently their roduct vanishes.



igure 22: Illustration of the piecewise linear basis functions corresponding to lobal node 2 and 3.

Salculating a specific matrix entry. Let us calculate the specific matrix atry $A_{2,3} = \int_{\Omega} \varphi_2 \varphi_3 dx$. Figure 22 shows how φ_2 and φ_3 look like. We realize om this figure that the product $\varphi_2 \varphi_3 \neq 0$ only over element 2, which contains

node 2 and 3. The particular formulas for $\varphi_2(x)$ and $\varphi_3(x)$ on $[x_2, x_3]$ ar from (54). The function φ_3 has positive slope over $[x_2, x_3]$ and corresp the interval $[x_{i-1}, x_i]$ in (54). With i = 3 we get

$$\varphi_3(x) = (x - x_2)/h,$$

while $\varphi_2(x)$ has negative slope over $[x_2, x_3]$ and corresponds to setting (54),

$$\varphi_2(x) = 1 - (x - x_2)/h$$
.

We can now easily integrate,

$$A_{2,3} = \int_{\Omega} \varphi_2 \varphi_3 \, dx = \int_{x_2}^{x_3} \left(1 - \frac{x - x_2}{h} \right) \frac{x - x_2}{h} \, dx = \frac{h}{6}.$$

The diagonal entry in the coefficient matrix becomes

$$A_{2,2} = \int_{x_1}^{x_2} \left(\frac{x - x_1}{h}\right)^2 dx + \int_{x_2}^{x_3} \left(1 - \frac{x - x_2}{h}\right)^2 dx = \frac{2h}{3}.$$

The entry $A_{2,1}$ has an the integral that is geometrically similar to the s in Figure 22, so we get $A_{2,1} = h/6$.

Calculating a general row in the matrix. We can now general calculation of matrix entries to a general row number i. The entry $A = \int_{\Omega} \varphi_i \varphi_{i-1} \, \mathrm{d}x$ involves hat functions as depicted in Figure 23. Since the is geometrically identical to the situation with specific nodes 2 and 3, we that $A_{i,i-1} = A_{i,i+1} = h/6$ and $A_{i,i} = 2h/3$. However, we can compute integral directly too:

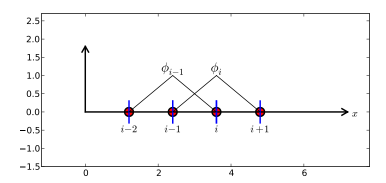
$$A_{i,i-1} = \int_{\Omega} \varphi_i \varphi_{i-1} \, \mathrm{d}x$$

$$= \underbrace{\int_{x_{i-2}}^{x_{i-1}} \varphi_i \varphi_{i-1} \, \mathrm{d}x}_{\varphi_i = 0} + \underbrace{\int_{x_{i-1}}^{x_i} \varphi_i \varphi_{i-1} \, \mathrm{d}x}_{\varphi_{i-1} = 0} + \underbrace{\int_{x_{i-1}}^{x_{i+1}} \varphi_i \varphi_{i-1} \, \mathrm{d}x}_{\varphi_{i-1} = 0}$$

$$= \underbrace{\int_{x_{i-1}}^{x_i} \underbrace{\left(\frac{x - x_i}{h}\right)}_{\varphi_i(x)} \underbrace{\left(1 - \frac{x - x_{i-1}}{h}\right)}_{\varphi_{i-1}(x)} \, \mathrm{d}x}_{\varphi_{i-1}(x)} = \frac{h}{6}.$$

The particular formulas for $\varphi_{i-1}(x)$ and $\varphi_i(x)$ on $[x_{i-1}, x_i]$ are four (54): φ_i is the linear function with positive slope, corresponding to the $[x_{i-1}, x_i]$ in (54), while ϕ_{i-1} has a negative slope so the definition in $[x_i, x_{i+1}]$ in (54) must be used. (The appearance of i in (54) and the might be confusing, as we speak about two different i indices.)

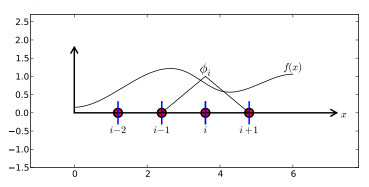
The first and last row of the coefficient matrix lead to slightly ϵ integrals:



igure 23: Illustration of two neighboring linear (hat) functions with general ode numbers.

$$A_{0,0} = \int_{\Omega} \varphi_0^2 dx = \int_{x_0}^{x_1} \left(1 - \frac{x - x_0}{h} \right)^2 dx = \frac{h}{3}.$$

imilarly, $A_{N,N}$ involves an integral over only one element and equals hence h/3.



igure 24: Right-hand side integral with the product of a basis function and ne given function to approximate.

The general formula for b_i , see Figure 24, is now easy to set up

$$f_i = \int_{\Omega} \varphi_i(x) f(x) \, \mathrm{d}x = \int_{x_{i-1}}^{x_i} \frac{x - x_{i-1}}{h} f(x) \, \mathrm{d}x + \int_{x_i}^{x_{i+1}} \left(1 - \frac{x - x_i}{h} \right) f(x) \, \mathrm{d}x.$$
(56)

/e need a specific f(x) function to compute these integrals. With two equal-sized ements in $\Omega = [0, 1]$ and f(x) = x(1 - x), one gets

$$A = \frac{h}{6} \begin{pmatrix} 2 & 1 & 0 \\ 1 & 4 & 1 \\ 0 & 1 & 2 \end{pmatrix}, \quad b = \frac{h^2}{12} \begin{pmatrix} 2 - 3h \\ 12 - 14h \\ 10 - 17h \end{pmatrix}.$$

The solution becomes

$$c_0 = \frac{h^2}{6}$$
, $c_1 = h - \frac{5}{6}h^2$, $c_2 = 2h - \frac{23}{6}h^2$.

The resulting function

$$u(x) = c_0 \varphi_0(x) + c_1 \varphi_1(x) + c_2 \varphi_2(x)$$

is displayed in Figure 25 (left). Doubling the number of elements to forto the improved approximation in the right part of Figure 25.

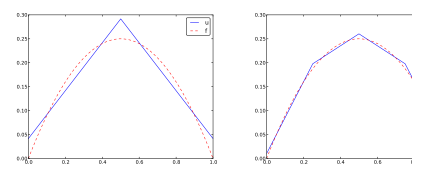


Figure 25: Least squares approximation of a parabola using 2 (left (right) P1 elements.

3.7 Assembly of elementwise computations

The integrals above are naturally split into integrals over individual e since the formulas change with the elements. This idea of splitting the is fundamental in all practical implementations of the finite element me Let us split the integral over Ω into a sum of contributions from each

$$A_{i,j} = \int_{\Omega} \varphi_i \varphi_j \, \mathrm{d}x = \sum_e A_{i,j}^{(e)}, \quad A_{i,j}^{(e)} = \int_{\Omega^{(e)}} \varphi_i \varphi_j \, \mathrm{d}x.$$

Now, $A_{i,j}^{(e)} \neq 0$ if and only if i and j are nodes in element e (look at Figurealize this property). Introduce i=q(e,r) as the mapping of local node r in element e to the global node number i. This is just a short mather notation for the expression i=elements[e][r] in a program. Let r a the local node numbers corresponding to the global node numbers i and j=q(e,s). With d nodes per element, all the nonzero elements in A from the integrals involving basis functions with indices corresponding global node numbers in element number e:

$$\int_{\Omega^{(e)}} \varphi_{q(e,r)} \varphi_{q(e,s)} \, \mathrm{d}x, \quad r, s = 0, \dots, d.$$

hese contributions can be collected in a $(d+1) \times (d+1)$ matrix known as the ement matrix. Let $I_d = \{0, \ldots, d\}$ be the valid indices of r and s. We introduce no notation

$$\tilde{A}^{(e)} = {\{\tilde{A}_{r,s}^{(e)}\}}, \quad r, s \in I_d,$$

or the element matrix. For the case d=2 we have

$$\tilde{A}^{(e)} = \begin{bmatrix} \tilde{A}_{0,0}^{(e)} & \tilde{A}_{0,1}^{(e)} & \tilde{A}_{0,2}^{(e)} \\ \tilde{A}_{1,0}^{(e)} & \tilde{A}_{1,1}^{(e)} & \tilde{A}_{1,2}^{(e)} \\ \tilde{A}_{2,0}^{(e)} & \tilde{A}_{2,1}^{(e)} & \tilde{A}_{2,2}^{(e)} \end{bmatrix}.$$

liven the numbers $\tilde{A}_{r,s}^{(e)}$, we should according to (57) add the contributions to ne global coefficient matrix by

$$A_{q(e,r),q(e,s)} := A_{q(e,r),q(e,s)} + \tilde{A}_{r,s}^{(e)}, \quad r,s \in I_d.$$
 (58)

his process of adding in elementwise contributions to the global matrix is called *nite element assembly* or simply *assembly*. Figure 26 illustrates how element natrices for elements with two nodes are added into the global matrix. More pecifically, the figure shows how the element matrix associated with elements 1 and 2 assembled, assuming that global nodes are numbered from left to right in 12 domain. With regularly numbered P3 elements, where the element matrices are size 4×4 , the assembly of elements 1 and 2 are sketched in Figure 27.

After assembly of element matrices corresponding to regularly numbered ements and nodes are understood, it is wise to study the assembly process for regularly numbered elements and nodes. Figure 17 shows a mesh where the lements array, or q(e,r) mapping in mathematical notation, is given as

he associated assembly of element matrices 1 and 2 is sketched in Figure 28.

These three assembly processes can also be animated⁶.

The right-hand side of the linear system is also computed elementwise:

$$b_i = \int_{\Omega} f(x)\varphi_i(x) dx = \sum_e b_i^{(e)}, \quad b_i^{(e)} = \int_{\Omega^{(e)}} f(x)\varphi_i(x) dx.$$
 (59)

We observe that $b_i^{(e)} \neq 0$ if and only if global node i is a node in element e (look to Figure 24 to realize this property). With d nodes per element we can collect the d+1 nonzero contributions $b_i^{(e)}$, for $i=q(e,r), r\in I_d$, in an element vector

$$\tilde{b}_r^{(e)} = \{\tilde{b}_r^{(e)}\}, \quad r \in I_d.$$

hese contributions are added to the global right-hand side by an assembly rocess similar to that for the element matrices:

$$b_{q(e,r)} := b_{q(e,r)} + \tilde{b}_r^{(e)}, \quad r \in I_d.$$
(60)

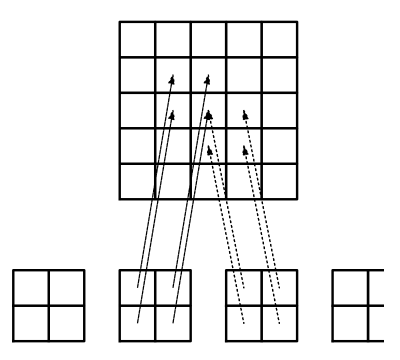


Figure 26: Illustration of matrix assembly: regularly numbered P1 ele

3.8 Mapping to a reference element

Instead of computing the integrals

$$\tilde{A}_{r,s}^{(e)} = \int_{\Omega^{(e)}} \varphi_{q(e,r)}(x) \varphi_{q(e,s)}(x) \, \mathrm{d}x$$

over some element $\Omega^{(e)} = [x_L, x_R]$, it is convenient to map the element $[x_L, x_R]$ to a standardized reference element domain [-1, 1]. (We have introduced x_L and x_R as the left and right boundary points of an avelement. With a natural, regular numbering of nodes and elements from right through the domain, we have $x_L = x_e$ and $x_R = x_{e+1}$ for P1 elements.

Let $X \in [-1, 1]$ be the coordinate in the reference element. A linear *apping* from X to x reads

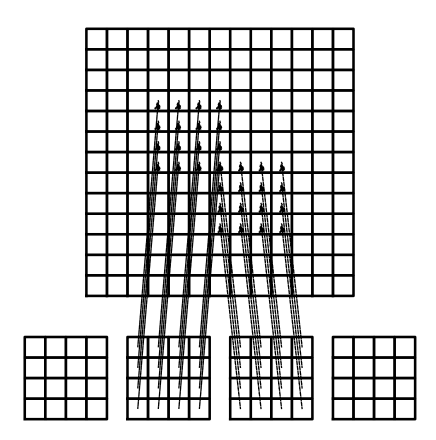
$$x = \frac{1}{2}(x_L + x_R) + \frac{1}{2}(x_R - x_L)X$$
.

This relation can alternatively be expressed by

$$x = x_m + \frac{1}{2}hX,$$

where we have introduced the element midpoint $x_m = (x_L + x_R)/2$ element length $h = x_R - x_L$.

⁶http://tinyurl.com/opdfafk/pub/mov-fem/fe_assembly.html



igure 27: Illustration of matrix assembly: regularly numbered P3 elements.

Integrating on the reference element is a matter of just changing the integraon variable from x to X. Let

$$\tilde{\varphi}_r(X) = \varphi_{q(e,r)}(x(X)) \tag{63}$$

e the basis function associated with local node number r in the reference ement. The integral transformation reads

$$\tilde{A}_{r,s}^{(e)} = \int_{\Omega^{(e)}} \varphi_{q(e,r)}(x) \varphi_{q(e,s)}(x) \, \mathrm{d}x = \int_{-1}^{1} \tilde{\varphi}_{r}(X) \tilde{\varphi}_{s}(X) \frac{dx}{dX} \, \mathrm{d}X. \tag{64}$$

he stretch factor dx/dX between the x and X coordinates becomes the determinant of the Jacobian matrix of the mapping between the coordinate systems 1 2D and 3D. To obtain a uniform notation for 1D, 2D, and 3D problems we herefore replace dx/dX by $\det J$ already now. In 1D, $\det J = dx/dX = h/2$, he integration over the reference element is then written as

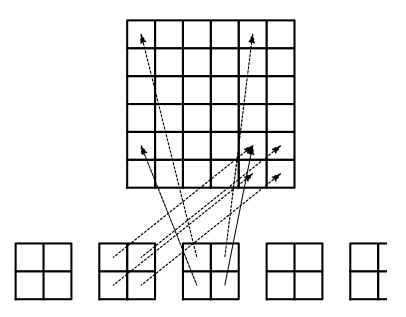


Figure 28: Illustration of matrix assembly: irregularly numbered P1 ele

$$\tilde{A}_{r,s}^{(e)} = \int_{-1}^{1} \tilde{\varphi}_r(X) \tilde{\varphi}_s(X) \det J \, dX \, .$$

The corresponding formula for the element vector entries becomes

$$\tilde{b}_r^{(e)} = \int_{\Omega^{(e)}} f(x) \varphi_{q(e,r)}(x) dx = \int_{-1}^1 f(x(X)) \tilde{\varphi}_r(X) \det J \, dX \,.$$

Since we from now on will work in the reference element, we need mathematical formulas for the basis functions $\varphi_i(x)$ in the reference only, i.e., we only need to specify formulas for $\tilde{\varphi}_r(X)$. This is a very consimplification compared to specifying piecewise polynomials in the planament.

The $\tilde{\varphi}_r(x)$ functions are simply the Lagrange polynomials defined the local nodes in the reference element. For d=1 and two nodes per we have the linear Lagrange polynomials

$$\tilde{\varphi}_0(X) = \frac{1}{2}(1 - X)$$

$$\tilde{\varphi}_1(X) = \frac{1}{2}(1 + X)$$

Quadratic polynomials, d = 2, have the formulas

$$\tilde{\varphi}_0(X) = \frac{1}{2}(X - 1)X$$
 (69)

$$\tilde{\varphi}_1(X) = 1 - X^2 \tag{70}$$

$$\tilde{\varphi}_2(X) = \frac{1}{2}(X+1)X\tag{71}$$

1 general.

$$\tilde{\varphi}_r(X) = \prod_{s=0, s \neq r}^d \frac{X - X_{(s)}}{X_{(r)} - X_{(s)}},\tag{72}$$

here $X_{(0)}, \ldots, X_{(d)}$ are the coordinates of the local nodes in the reference ement. These are normally uniformly spaced: $X_{(r)} = -1 + 2r/d$, $r \in I_d$.

Why reference elements?

The great advantage of using reference elements is that the formulas for the basis functions, $\tilde{\varphi}_r(X)$, are the same for all elements and independent of the element geometry (length and location in the mesh). The geometric information is "factored out" in the simple mapping formula and the associated det J quantity, but this information is (here taken as) the same for element types. Also, the integration domain is the same for all elements.

.9 Example: Integration over a reference element

o illustrate the concepts from the previous section in a specific example, we ow consider calculation of the element matrix and vector for a specific choice f d and f(x). A simple choice is d=1 (P1 elements) and f(x)=x(1-x) n $\Omega=[0,1]$. We have the general expressions (65) and (66) for $\tilde{A}_{r,s}^{(e)}$ and $\tilde{b}_r^{(e)}$. /riting these out for the choices (67) and (68), and using that $\det J=h/2$, we an do the following calculations of the element matrix entries:

$$\begin{split} \tilde{A}_{0,0}^{(e)} &= \int_{-1}^{1} \tilde{\varphi}_{0}(X) \tilde{\varphi}_{0}(X) \frac{h}{2} dX \\ &= \int_{-1}^{1} \frac{1}{2} (1-X) \frac{1}{2} (1-X) \frac{h}{2} dX = \frac{h}{8} \int_{-1}^{1} (1-X)^{2} dX = \frac{h}{3}, \\ \tilde{A}_{1,0}^{(e)} &= \int_{-1}^{1} \tilde{\varphi}_{1}(X) \tilde{\varphi}_{0}(X) \frac{h}{2} dX \\ &= \int_{-1}^{1} \frac{1}{2} (1+X) \frac{1}{2} (1-X) \frac{h}{2} dX = \frac{h}{8} \int_{-1}^{1} (1-X^{2}) dX = \frac{h}{6}, \\ \tilde{A}_{0,1}^{(e)} &= \tilde{A}_{1,0}^{(e)}, \\ \tilde{A}_{1,1}^{(e)} &= \int_{-1}^{1} \tilde{\varphi}_{1}(X) \tilde{\varphi}_{1}(X) \frac{h}{2} dX \\ &= \int_{-1}^{1} \frac{1}{2} (1+X) \frac{1}{2} (1+X) \frac{h}{2} dX = \frac{h}{8} \int_{-1}^{1} (1+X)^{2} dX = \frac{h}{3} \,. \end{split}$$

The corresponding entries in the element vector becomes

$$\begin{split} \tilde{b}_0^{(e)} &= \int_{-1}^1 f(x(X)) \tilde{\varphi}_0(X) \frac{h}{2} dX \\ &= \int_{-1}^1 (x_m + \frac{1}{2} hX) (1 - (x_m + \frac{1}{2} hX)) \frac{1}{2} (1 - X) \frac{h}{2} dX \\ &= -\frac{1}{24} h^3 + \frac{1}{6} h^2 x_m - \frac{1}{12} h^2 - \frac{1}{2} h x_m^2 + \frac{1}{2} h x_m \\ \tilde{b}_1^{(e)} &= \int_{-1}^1 f(x(X)) \tilde{\varphi}_1(X) \frac{h}{2} dX \\ &= \int_{-1}^1 (x_m + \frac{1}{2} hX) (1 - (x_m + \frac{1}{2} hX)) \frac{1}{2} (1 + X) \frac{h}{2} dX \\ &= -\frac{1}{24} h^3 - \frac{1}{6} h^2 x_m + \frac{1}{12} h^2 - \frac{1}{2} h x_m^2 + \frac{1}{2} h x_m \,. \end{split}$$

In the last two expressions we have used the element midpoint x_m .

Integration of lower-degree polynomials above is tedious, and higher polynomials involve very much more algebra, but sympy may help. For ϵ we can easily calculate (73), (73), and (77) by

```
>>> import sympy as sp
>>> x, x_m, h, X = sp.symbols('x x_m h X')
>>> sp.integrate(h/8*(1-X)**2, (X, -1, 1))
h/3
>>> sp.integrate(h/8*(1+X)*(1-X), (X, -1, 1))
h/6
>>> x = x_m + h/2*X
>>> b_0 = sp.integrate(h/4*x*(1-x)*(1-X), (X, -1, 1))
```

```
>>> print b_0
-h**3/24 + h**2*x_m/6 - h**2/12 - h*x_m**2/2 + h*x_m/2
```

or inclusion of formulas in documents (like the present one), sympy can print spressions in IATEX format:

```
>>> print sp.latex(b_0, mode='plain')
- \frac{1}{24} h^{3} + \frac{1}{6} h^{2} x_{m}
- \frac{1}{12} h^{2} - \half h x_{m}^{2}
+ \half h x_{m}
```

Implementation

ased on the experience from the previous example, it makes sense to write ome code to automate the analytical integration process for any choice of finite lement basis functions. In addition, we can automate the assembly process nd linear system solution. Appropriate functions for this purpose document ll details of all steps in the finite element computations and can found in the rodule file fe_approx1D.py⁷. The key steps in the computational machinery re now explained in detail in terms of code and text.

.1 Integration

irst we need a Python function for defining $\tilde{\varphi}_r(X)$ in terms of a Lagrange olynomial of degree d:

```
import sympy as sp
import numpy as np
lef phi_r(r, X, d):
   if isinstance(X, sp.Symbol):
       h = sp.Rational(1, d) # node spacing
       nodes = [2*i*h - 1 \text{ for } i \text{ in } range(d+1)]
   else:
       # assume X is numeric: use floats for nodes
       nodes = np.linspace(-1, 1, d+1)
   return Lagrange_polynomial(X, r, nodes)
lef Lagrange_polynomial(x, i, points):
   p = 1
   for k in range(len(points)):
       if k != i:
           p *= (x - points[k])/(points[i] - points[k])
   return p
```

bserve how we construct the phi_r function to be a symbolic expression for r(X) if X is a Symbol object from sympy. Otherwise, we assume that X is a float bject and compute the corresponding floating-point value of $\tilde{\varphi}_r(X)$. Recall that

the Lagrange_polynomial function, here simply copied from Section 2.' with both symbolic and numeric variables.

The complete basis $\tilde{\varphi}_0(X), \ldots, \tilde{\varphi}_d(X)$ on the reference element, repr as a list of symbolic expressions, is constructed by

```
def basis(d=1):
    X = sp.Symbol('X')
    phi = [phi_r(r, X, d) for r in range(d+1)]
    return phi
```

Now we are in a position to write the function for computing the element

In the symbolic case (symbolic is True), we introduce the element le a symbol h in the computations. Otherwise, the real numerical value element interval Omega_e is used and the final matrix elements are numb symbols. This functionality can be demonstrated:

The computation of the element vector is done by a similar procedu

```
def element_vector(f, phi, Omega_e, symbolic=True):
    n = len(phi)
    b_e = sp.zeros((n, 1))
    # Make f a function of X
    X = sp.Symbol('X')
    if symbolic:
        h = sp.Symbol('h')
    else:
        h = Omega_e[1] - Omega_e[0]
    x = (Omega_e[0] + Omega_e[1])/2 + h/2*X # mapping
```

⁷http://tinyurl.com/nm5587k/fem/fe_approx1D.py

```
f = f.subs('x', x) # substitute mapping formula for x
detJ = h/2 # dx/dX
for r in range(n):
    b_e[r] = sp.integrate(f*phi[r]*detJ, (X, -1, 1))
return b_e
```

lere we need to replace the symbol \mathbf{x} in the expression for \mathbf{f} by the mapping rmula such that \mathbf{f} can be integrated in terms of X, cf. the formula $\tilde{b}_r^{(e)} = \frac{1}{-1} f(x(X)) \tilde{\varphi}_r(X) \frac{h}{2} dX$.

The integration in the element matrix function involves only products of olynomials, which sympy can easily deal with, but for the right-hand side sympy ay face difficulties with certain types of expressions f. The result of the integral then an Integral object and not a number or expression as when symbolic itegration is successful. It may therefore be wise to introduce a fallback on umerical integration. The symbolic integration can also take much time before a unsuccessful conclusion so we may also introduce a parameter symbolic and it to False in order to avoid symbolic integration:

```
lef element_vector(f, phi, Omega_e, symbolic=True):
    ...
    if symbolic:
        I = sp.integrate(f*phi[r]*detJ, (X, -1, 1))
    if not symbolic or isinstance(I, sp.Integral):
        h = Omega_e[1] - Omega_e[0] # Ensure h is numerical
        detJ = h/2
        integrand = sp.lambdify([X], f*phi[r]*detJ)
        I = sp.mpmath.quad(integrand, [-1, 1])
        b_e[r] = I
    ...
```

umerical integration requires that the symbolic integrand is converted to a plain ython function (integrand) and that the element length h is a real number.

.2 Linear system assembly and solution

he complete algorithm for computing and assembling the elementwise contribuous takes the following form

```
lef assemble(nodes, elements, phi, f, symbolic=True):
    N_n, N_e = len(nodes), len(elements)
    if symbolic:
        A = sp.zeros((N_n, N_n))
        b = sp.zeros((N_n, 1))  # note: (N_n, 1) matrix
    else:
        A = np.zeros((N_n, N_n))
        b = np.zeros(N_n)
    for e in range(N_e):
        Omega_e = [nodes[elements[e][0]], nodes[elements[e][-1]]]

        A_e = element_matrix(phi, Omega_e, symbolic)
        b_e = element_vector(f, phi, Omega_e, symbolic)
        for r in range(len(elements[e])):
```

The nodes and elements variables represent the finite element mesh as exearlier

Given the coefficient matrix A and the right-hand side b, we can c the coefficients $\{c_i\}_{i\in\mathcal{I}_s}$ in the expansion $u(x)=\sum_j c_j\varphi_j$ as the solution c of the linear system:

```
if symbolic:
    c = A.LUsolve(b)
else:
    c = np.linalg.solve(A, b)
```

When A and b are sympy arrays, the solution procedure implied by A.LUs symbolic. Otherwise, A and b are numpy arrays and a standard numerical is called. The symbolic version is suited for small problems only (small A since the calculation time becomes prohibitively large otherwise. Normal symbolic integration will be more time consuming in small problems to symbolic solution of the linear system.

4.3 Example on computing symbolic approximation

We can exemplify the use of assemble on the computational case from Sec with two P1 elements (linear basis functions) on the domain $\Omega = [0, 1]$ first work with a symbolic element length:

```
>>> h, x = sp.symbols('h x')
>>> nodes = [0, h, 2*h]
>>> elements = [[0, 1], [1, 2]]
>>> phi = basis(d=1)
>>> f = x*(1-x)
>>> A, b = assemble(nodes, elements, phi, f, symbolic=True)
[h/3, h/6, 0]
[h/6, 2*h/3, h/6]
[0, h/6, h/3]
>>> b
     h**2/6 - h**3/12]
      h**2 - 7*h**3/6]
[5*h**2/6 - 17*h**3/12]
>>> c = A.LUsolve(b)
>>> c
                           h**2/6]
[12*(7*h**2/12 - 35*h**3/72)/(7*h)]
[7*(4*h**2/7 - 23*h**3/21)/(2*h)]
```

.4 Comparison with finite elements and interpolation/collocation

/e may, for comparison, compute the ${\tt c}$ vector corresponding to an interpolaon/collocation method with finite element basis functions. Choosing the nodes ${\tt s}$ points, the principle is

$$u(x_i) = \sum_{j \in \mathcal{I}_s} c_j \varphi_j(x_i) = f(x_i), \quad i \in \mathcal{I}_s.$$

he coefficient matrix $A_{i,j} = \varphi_j(x_i)$ becomes the identity matrix because basis unction number j vanishes at all nodes, except node j: $\varphi_j(x_i = \delta_{ij})$. Therefore, $j = f(x_i)$.

The associated sympy calculations are

```
>>> fn = sp.lambdify([x], f)
>>> c = [fn(xc) for xc in nodes]
>>> c
[0, h*(1 - h), 2*h*(1 - 2*h)]
```

hese expressions are much simpler than those based on least squares or projecon in combination with finite element basis functions.

5 Example on computing numerical approximations

he numerical computations corresponding to the symbolic ones in Section 4.3, and still done by sympy and the assemble function, go as follows:

```
\rightarrow nodes = [0, 0.5, 1]
>>> elements = [[0, 1], [1, 2]]
>>> phi = basis(d=1)
>>> x = sp.Symbol('x')
\Rightarrow f = x*(1-x)
>>> A, b = assemble(nodes, elements, phi, f, symbolic=False)
[0.083333333333333, 0.3333333333333, 0.0833333333333333]
               >>> h
         0.03125]
[0.104166666666667]
         0.031257
>>> c = A.LUsolve(b)
>>> c
[0.041666666666666]
 0.2916666666666671
[0.041666666666666]
```

The fe_approx1D module contains functions for generating the nodes and lements lists for equal-sized elements with any number of nodes per element. he coordinates in nodes can be expressed either through the element length /mbol h (symbolic=True) or by real numbers (symbolic=False):

There is also a function

```
def approximate(f, symbolic=False, d=1, N_e=4, filename='tmp.pdf'
```

which computes a mesh with N_e elements, basis functions of degree approximates a given symbolic expression f by a finite element expansion $\sum_j c_j \varphi_j(x)$. When symbolic is False, $u(x) = \sum_j c_j \varphi_j(x)$ can be compa (large) number of points and plotted together with f(x). The construct points from the solution vector c is done elementwise by evaluating $\sum_r c_r (a_r + b_r) = 0$ number of points in each element in the local coordinate and the discrete (x, u) values on each element are stored in separate arrare finally concatenated to form a global array for x and for x. The defound in the u_glob function in fe_approx1D.py.

4.6 The structure of the coefficient matrix

Let us first see how the global matrix looks like if we assemble symbolic matrices, expressed in terms of h, from several elements:

```
>>> d=1; N_e=8; Omega=[0,1] # 8 linear elements on [0,1]
>>> phi = basis(d)
>> f = x*(1-x)
>>> nodes, elements = mesh_symbolic(N_e, d, Omega)
>>> A. b = assemble(nodes, elements, phi, f, symbolic=True)
>>> A
[h/3, h/6,
[h/6, 2*h/3,
            h/6.
                    0.
                            0.
                                                     01
       h/6, 2*h/3, h/6,
                           0,
                                                     0]
              h/6, 2*h/3, h/6,
                                                     01
             0, h/6, 2*h/3, h/6,
                                          0,
[ 0,
[ 0,
                    0,
                          h/6, 2*h/3,
                                        h/6,
         0,
               0,
[ 0,
         0,
             0,
                      0,
                             0,
                                 h/6, 2*h/3,
                                               h/6,
[ 0,
         0,
                      0,
                                   0,
                                        h/6, 2*h/3, h/6]
               0,
                             0,
[ 0,
         0.
               0.
                      0,
                                   0,
                                          0.
                                               h/6, h/3
```

The reader is encouraged to assemble the element matrices by hand an this result, as this exercise will give a hands-on understanding of w assembly is about. In general we have a coefficient matrix that is tridia

The structure of the right-hand side is more difficult to reveal since it involves n assembly of elementwise integrals of $f(x(X))\tilde{\varphi}_r(X)h/2$, which obviously epend on the particular choice of f(x). Numerical integration can give some sight into the nature of the right-hand side. For this purpose it is easier to bok at the integration in x coordinates, which gives the general formula (56). or equal-sized elements of length h, we can apply the Trapezoidal rule at the lobal node points to arrive at

$$b_{i} = h \left(\frac{1}{2} \varphi_{i}(x_{0}) f(x_{0}) + \frac{1}{2} \varphi_{i}(x_{N}) f(x_{N}) + \sum_{j=1}^{N-1} \varphi_{i}(x_{j}) f(x_{j}) \right)$$
(80)

$$= \begin{cases} \frac{1}{2}hf(x_i), & i = 0 \text{ or } i = N, \\ hf(x_i), & 1 \le i \le N - 1 \end{cases}$$
 (81)

he reason for this simple formula is simply that φ_i is either 0 or 1 at the nodes and 0 at all but one of them.

Going to P2 elements (d=2) leads to the element matrix

$$A^{(e)} = \frac{h}{30} \begin{pmatrix} 4 & 2 & -1\\ 2 & 16 & 2\\ -1 & 2 & 4 \end{pmatrix}$$
 (82)

nd the following global assembled matrix from four elements:

$$A = \frac{h}{30} \begin{pmatrix} 4 & 2 & -1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 2 & 16 & 2 & 0 & 0 & 0 & 0 & 0 & 0 \\ -1 & 2 & 8 & 2 & -1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 2 & 16 & 2 & 0 & 0 & 0 & 0 \\ 0 & 0 & -1 & 2 & 8 & 2 & -1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 2 & 16 & 2 & 0 & 0 \\ 0 & 0 & 0 & 0 & -1 & 2 & 8 & 2 & -1 \\ 0 & 0 & 0 & 0 & 0 & 0 & 2 & 16 & 2 \\ 0 & 0 & 0 & 0 & 0 & 0 & 2 & 16 & 2 \\ 0 & 0 & 0 & 0 & 0 & 0 & -1 & 2 & 4 \end{pmatrix}$$
 (83)

In general, for i odd we have the nonzeroes

$$A_{i,i-2} = -1$$
, $A_{i-1,i} = 2$, $A_{i,i} = 8$, $A_{i+1,i} = 2$, $A_{i+2,i} = -1$

multiplied by h/30, and for i even we have the nonzeros

$$A_{i-1,i} = 2$$
, $A_{i,i} = 16$, $A_{i+1,i} = 2$,

multiplied by h/30. The rows with odd numbers correspond to node element boundaries and get contributions from two neighboring element assembly process, while the even numbered rows correspond to internal 1 the elements where the only one element contributes to the values in th matrix.

4.7 Applications

With the aid of the approximate function in the fe_approx1D module easily investigate the quality of various finite element approximations given functions. Figure 29 shows how linear and quadratic elements approximate polynomial $f(x) = x(1-x)^8$ on $\Omega = [0,1]$, using equal-sized element results arise from the program

```
import sympy as sp
from fe_approx1D import approximate
x = sp.Symbol('x')

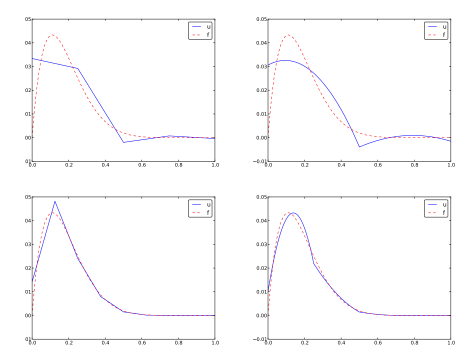
approximate(f=x*(1-x)**8, symbolic=False, d=1, N_e=4)
approximate(f=x*(1-x)**8, symbolic=False, d=2, N_e=2)
approximate(f=x*(1-x)**8, symbolic=False, d=1, N_e=8)
approximate(f=x*(1-x)**8, symbolic=False, d=2, N_e=4)
```

The quadratic functions are seen to be better than the linear ones for the value of N, as we increase N. This observation has some generality: degree is not necessarily better on a coarse mesh, but it is as we refilt mesh.

4.8 Sparse matrix storage and solution

Some of the examples in the preceding section took several minutes to c even on small meshes consisting of up to eight elements. The main expl for slow computations is unsuccessful symbolic integration: sympy malot of energy on integrals like $\int f(x(X))\tilde{\varphi}_r(X)h/2dx$ before giving up, program then resorts to numerical integration. Codes that can deal with number of basis functions and accept flexible choices of f(x) should com integrals numerically and replace the matrix objects from sympy by the f efficient array objects from numpy.

Another reason for slow code is related to the fact that most of the entries $A_{i,j}$ are zero, because $(\varphi_i, \varphi_j) = 0$ unless i and j are nodes in the



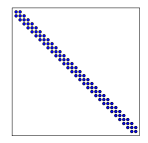
igure 29: Comparison of the finite element approximations: 4 P1 elements with nodes (upper left), 2 P2 elements with 5 nodes (upper right), 8 P1 elements ith 9 nodes (lower left), and 4 P2 elements with 9 nodes (lower right).

lement. A matrix whose majority of entries are zeros, is known as a *sparse* latrix. The sparsity should be utilized in software as it dramatically decreases the storage demands and the CPU-time needed to compute the solution of the near system. This optimization is not critical in 1D problems where modern omputers can afford computing with all the zeros in the complete square matrix, ut in 2D and especially in 3D, sparse matrices are fundamental for feasible nite element computations.

In 1D problems, using a numbering of nodes and elements from left to right ver the domain, the assembled coefficient matrix has only a few diagonals ifferent from zero. More precisely, 2d+1 diagonals are different from zero. 7 ith a different numbering of global nodes, say a random ordering, the diagonal ructure is lost, but the number of nonzero elements is unaltered. Figures 30 and 31 exemplify sparsity patterns.

The scipy.sparse library supports creation of sparse matrices and linear stem solution.

- \bullet scipy.sparse.diags for matrix defined via diagonals
- scipy.sparse.lil_matrix for creation via setting matrix entries



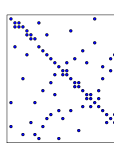
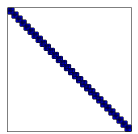


Figure 30: Matrix sparsity pattern for left-to-right numbering (left) and numbering (right) of nodes in P1 elements.



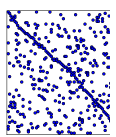


Figure 31: Matrix sparsity pattern for left-to-right numbering (left) and numbering (right) of nodes in P3 elements.

• scipy.sparse.dok_matrix for creation via setting matrix entries

5 Comparison of finite element and finite c ence approximation

The previous sections on approximating f by a finite element function of the projection/Galerkin or least squares approaches to minimize the amation error. We may, alternatively, use the collocation/interpolation as described in Section 4.4. Here we shall compare these three approach what one does in the finite difference method when representing a given to on a mesh.

5.1 Finite difference approximation of given function

Approximating a given function f(x) on a mesh in a finite difference cont typically just sample f at the mesh points. If u_i is the value of the approx at the mesh point x_i , we have $u_i = f(x_i)$. The collocation/interpolation using finite element basis functions gives exactly the same representa shown Section 4.4,

$$u(x_i) = c_i = f(x_i).$$

How does a finite element Galerkin or least squares approximation differ from its straightforward interpolation of f? This is the question to be addressed ext. We now limit the scope to P1 elements since this is the element type that ives formulas closest to those arising in the finite difference method.

.2 Finite difference interpretation of a finite element approximation

he linear system arising from a Galerkin or least squares approximation reads ι general

$$\sum_{j \in \mathcal{I}_s} c_j(\psi_i, \psi_j) = (f, \psi_i), \quad i \in \mathcal{I}_s.$$

1 the finite element approximation we choose $\psi_i = \varphi_i$. With φ_i corresponding 1 P1 elements and a uniform mesh of element length h we have in Section 3.6 alculated the matrix with entries (φ_i, φ_j) . Equation number i reads

$$\frac{h}{6}(u_{i-1} + 4u_i + u_{i+1}) = (f, \varphi_i). \tag{84}$$

he first and last equation, corresponding to i=0 and i=N are slightly ifferent, see Section 4.6.

The finite difference counterpart to (84) is just $u_i = f_i$ as explained in ection 5.1. To easier compare this result to the finite element approach to pproximating functions, we can rewrite the left-hand side of (84) as

$$h(u_i + \frac{1}{6}(u_{i-1} - 2u_i + u_{i+1})). (85)$$

hinking in terms of finite differences, we can write this expression using finite ifference operator notation:

$$[h(u+\frac{h^2}{6}D_xD_xu)]_i,$$

hich is nothing but the standard discretization of

$$h(u+\frac{h^2}{6}u'')$$
.

Before interpreting the approximation procedure as solving a differential quation, we need to work out what the right-hand side is in the context of P1 lements. Since φ_i is the linear function that is 1 at x_i and zero at all other odes, only the interval $[x_{i-1}, x_{i+1}]$ contribute to the integral on the right-hand de. This integral is naturally split into two parts according to (54):

$$(f,\varphi_i) = \int_{x_{i-1}}^{x_i} f(x) \frac{1}{h} (x - x_{i-1}) dx + \int_{x_i}^{x_{i+1}} f(x) \frac{1}{h} (1 - (x - x_i)) dx$$

However, if f is not known we cannot do much else with this expressic clear that many values of f around x_i contributes to the right-hand s just the single point value $f(x_i)$ as in the finite difference method.

To proceed with the right-hand side, we can turn to numerical inteschemes. The Trapezoidal method for (f, φ_i) , based on sampling the in $f\varphi_i$ at the node points $x_i = ih$ gives

$$(f,\varphi_i) = \int_{\Omega} f\varphi_i dx \approx h \frac{1}{2} (f(x_0)\varphi_i(x_0) + f(x_N)\varphi_i(x_N)) + h \sum_{j=1}^{N-1} f(x_j)\varphi_j(x_N) + h \sum_{j=1}^{N-1} f(x_N)\varphi_j(x_N) + h \sum_{j=1}^{N-1} f(x_N)\varphi_j$$

Since φ_i is zero at all these points, except at x_i , the Trapezoidal rule c to one term:

$$(f, \varphi_i) \approx hf(x_i),$$

for i = 1, ..., N - 1, which is the same result as with collocation/interprand of course the same result as in the finite difference method. For i : i = N we get contribution from only one element so

$$(f, \varphi_i) \approx \frac{1}{2} h f(x_i), \quad i = 0, \ i = N.$$

Simpson's rule with sample points also in the middle of the elem $x_{i+\frac{1}{2}} = (x_i + x_{i+1})/2$, can be written as

$$\int_{\Omega} g(x)dx \approx \frac{\tilde{h}}{3} \left(g(x_0) + 2 \sum_{j=1}^{N-1} g(x_j) + 4 \sum_{j=0}^{N-1} g(x_{j+\frac{1}{2}}) + f(x_{2N}) \right)$$

where $\tilde{h} = h/2$ is the spacing between the sample points. Our integrand $f\varphi_i$. For all the node points, $\varphi_i(x_j) = \delta_{ij}$, and therefore $\sum_{j=1}^{N-1} f(x_j) \varphi_j$ $f(x_i)$. At the midpoints, $\varphi_i(x_{i\pm\frac{1}{2}}) = 1/2$ and $\varphi_i(x_{j+\frac{1}{2}}) = 0$ for j > j < i-1. Consequently,

$$\sum_{j=0}^{N-1} f(x_{j+\frac{1}{2}}) \varphi_i(x_{j+\frac{1}{2}}) = \frac{1}{2} (fx_{j-\frac{1}{2}} + x_{j+\frac{1}{2}}) \,.$$

When $1 \le i \le N-1$ we then get

$$(f, \varphi_i) \approx \frac{h}{3} (f_{i-\frac{1}{2}} + f_i + f_{i+\frac{1}{2}}).$$

his result shows that, with Simpson's rule, the finite element method operates ith the average of f over three points, while the finite difference method just pplies f at one point. We may interpret this as a "smearing" or smoothing of f y the finite element method.

We can now summarize our findings. With the approximation of (f, φ_i) by ne Trapezoidal rule, P1 elements give rise to equations that can be expressed a finite difference discretization of

$$u + \frac{h^2}{6}u'' = f, \quad u'(0) = u'(L) = 0,$$
 (89)

spressed with operator notation as

$$[u + \frac{h^2}{6} D_x D_x u = f]_i. (90)$$

s $h \to 0$, the extra term proportional to u'' goes to zero, and the two methods re then equal.

With the Simpson's rule, we may say that we solve

$$[u + \frac{h^2}{6} D_x D_x u = \bar{f}]_i, (91)$$

here \bar{f}_i means the average $\frac{1}{3}(f_{i-1/2}+f_i+f_{i+1/2})$.

The extra term $\frac{h^2}{6}u''$ represents a smoothing effect: with just this term, e would find u by integrating f twice and thereby smooth f considerably. In addition, the finite element representation of f involves an average, or a moothing, of f on the right-hand side of the equation system. If f is a noisy metion, direct interpolation $u_i = f_i$ may result in a noisy u too, but with a alerkin or least squares formulation and P1 elements, we should expect that u smoother than f unless h is very small.

The interpretation that finite elements tend to smooth the solution is valid applications far beyond approximation of 1D functions.

3 Making finite elements behave as finite differences

/ith a simple trick, using numerical integration, we can easily produce the result $i = f_i$ with the Galerkin or least square formulation with P1 elements. This is seful in many occasions when we deal with more difficult differential equations and want the finite element method to have properties like the finite difference nethod (solving standard linear wave equations is one primary example).

Computations in physical space. We have already seen that applying the rapezoidal rule to the right-hand side (f, φ_i) simply gives f sampled at x_i . sing the Trapezoidal rule on the matrix entries $A_{i,j} = (\varphi_i, \varphi_j)$ involves a sum

$$\sum_{k} \varphi_i(x_k) \varphi_j(x_k),$$

but $\varphi_i(x_k) = \delta_{ik}$ and $\varphi_j(x_k) = \delta_{jk}$. The product $\varphi_i\varphi_j$ is then differe zero only when sampled at x_i and i = j. The Trapezoidal approximation integral is then

$$(\varphi_i, \varphi_j) \approx h, \quad i = j,$$

and zero if $i \neq j$. This means that we have obtained a diagonal matrix! and last diagonal elements, (φ_0, φ_0) and (φ_N, φ_N) get contribution on the first and last element, respectively, resulting in the approximate value h/2. The corresponding right-hand side also has a factor 1/2 for i = N. Therefore, the least squares or Galerkin approach with P1 elemetrapezoidal integration results in

$$c_i = f_i, \quad i \in \mathcal{I}_s$$
.

Simpsons's rule can be used to achieve a similar result for P2 elementiagonal coefficient matrix, but with the previously derived average of jurisht-hand side.

Elementwise computations. Identical results to those above will we perform elementwise computations. The idea is to use the Trapezoi on the reference element for computing the element matrix and vector assembled, the same equations $c_i = f(x_i)$ arise. Exercise 19 encourages carry out the details.

Terminology. The matrix with entries (φ_i, φ_j) typically arises from proportional to u in a differential equation where u is the unknown f This matrix is often called the *mass matrix*, because in the early day finite element method, the matrix arose from the mass times accelerated in Newton's second law of motion. Making the mass matrix diagonal numerical integration, as demonstrated above, is a widely used technique called *mass lumping*. In time-dependent problems it can sometimes enhancement accuracy and computational efficiency of the finite element. However, there are also examples where mass lumping destroys accuracy

6 A generalized element concept

So far, finite element computing has employed the nodes and element together with the definition of the basis functions in the reference of Suppose we want to introduce a piecewise constant approximation with of function $\tilde{\varphi}_0(x) = 1$ in the reference element, corresponding to a $\varphi_i(x)$ function in the element number i and zero on all other elements. Although i associate the function value with a node in the middle of the elements, to no nodes at the ends, and the previous code snippets will not work because the find the element boundaries from the nodes list.

.1 Cells, vertices, and degrees of freedom

/e now introduce cells as the subdomains $\Omega^{(e)}$ previously referred as elements. he cell boundaries are denoted as vertices. The reason for this name is that ells are recognized by their vertices in 2D and 3D. We also define a set of degrees freedom, which are the quantities we aim to compute. The most common type f degree of freedom is the value of the unknown function u at some point. (For cample, we can introduce nodes as before and say the degrees of freedom are the alues of u at the nodes.) The basis functions are constructed so that they equal nity for one particular degree of freedom and zero for the rest. This property names that when we evaluate $u = \sum_j c_j \varphi_j$ for degree of freedom number i, we et $u = c_i$. Integrals are performed over cells, usually by mapping the cell of iterest to a reference cell.

With the concepts of cells, vertices, and degrees of freedom we increase the ecoupling of the geometry (cell, vertices) from the space of basis functions. We will associate different sets of basis functions with a cell. In 1D, all cells re intervals, while in 2D we can have cells that are triangles with straight des, or any polygon, or in fact any two-dimensional geometry. Triangles and uadrilaterals are most common, though. The popular cell types in 3D are strahedra and hexahedra.

.2 Extended finite element concept

he concept of a finite element is now

- a reference cell in a local reference coordinate system;
- a set of basis functions $\tilde{\varphi}_i$ defined on the cell;
- a set of degrees of freedom that uniquely determines the basis functions such that $\tilde{\varphi}_i = 1$ for degree of freedom number i and $\tilde{\varphi}_i = 0$ for all other degrees of freedom;
- a mapping between local and global degree of freedom numbers, here called the *dof map*;
- a geometric mapping of the reference cell onto to cell in the physical domain.

here must be a geometric description of a cell. This is trivial in 1D since the ell is an interval and is described by the interval limits, here called vertices. If ne cell is $\Omega^{(e)} = [x_L, x_R]$, vertex 0 is x_L and vertex 1 is x_R . The reference cell 1D is [-1, 1] in the reference coordinate system X.

The expansion of u over one cell is often used:

$$u(x) = \tilde{u}(X) = \sum_{r} c_r \tilde{\varphi}_r(X), \quad x \in \Omega^{(e)}, \ X \in [-1, 1],$$
 (92)

where the sum is taken over the numbers of the degrees of freedom and ϵ value of u for degree of freedom number r.

Our previous P1, P2, etc., elements are defined by introducing d+1 spaced nodes in the reference cell and saying that the degrees of freedom d+1 function values at these nodes. The basis functions must be 1 at 0 and 0 at the others, and the Lagrange polynomials have exactly this p. The nodes can be numbered from left to right with associated degrees of that are numbered in the same way. The degree of freedom mapping $\mathfrak k$ what was previously represented by the elements lists. The cell mappir same affine mapping (61) as before.

6.3 Implementation

Implementationwise,

- we replace nodes by vertices;
- we introduce cells such that cell[e][r] gives the mapping frovertex r in cell e to the global vertex number in vertices;
- we replace elements by dof_map (the contents are the same for ments).

Consider the example from Section 3.1 where $\Omega = [0,1]$ is divided into t $\Omega^{(0)} = [0,0.4]$ and $\Omega^{(1)} = [0.4,1]$, as depicted in Figure 16. The vert [0,0.4,1]. Local vertex 0 and 1 are 0 and 0.4 in cell 0 and 0.4 and 1 in α P2 element means that the degrees of freedom are the value of α at three spaced points (nodes) in each cell. The data structures become

```
vertices = [0, 0.4, 1]
cells = [[0, 1], [1, 2]]
dof_map = [[0, 1, 2], [2, 3, 4]]
```

If we would approximate f by piecewise constants, known as P0 elem simply introduce one point or node in an element, preferably X=0, an one degree of freedom, which is the function value at this node. Moreover, set $\tilde{\varphi}_0(X)=1$. The cells and vertices arrays remain the same, but $\tilde{\varphi}_0(X)=1$ is altered:

```
dof_map = [[0], [1]]
```

We use the cells and vertices lists to retrieve information on the g of a cell, while dof_map is the q(e,r) mapping introduced earlier in the a of element matrices and vectors. For example, the Omega_e variable (reprethe cell interval) in previous code snippets must now be computed as

Dmega e = [vertices[cells[e][0], vertices[cells[e][1]]

he assembly is done by

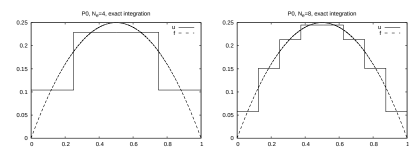
```
\[dof_map[e][r], dof_map[e][s]] += A_e[r,s]
>[dof_map[e][r]] += b_e[r]
```

We will hereafter drop the nodes and elements arrays and work exclusively ith cells, vertices, and dof_map. The module fe_approx1D_numint.py ow replaces the module fe_approx1D and offers similar functions that work ith the new concepts:

hese steps are offered in the approximate function, which we here apply to see ow well four P0 elements (piecewise constants) can approximate a parabola:

```
from fe_approx1D_numint import *
c=sp.Symbol("x")
for N_e in 4, 8:
    approximate(x*(1-x), d=0, N_e=N_e, Omega=[0,1])
```

igure 32 shows the result.



igure 32: Approximation of a parabola by 4 (left) and 8 (right) P0 elements.

6.4 Computing the error of the approximation

So far we have focused on computing the coefficients c_j in the approx $u(x) = \sum_j c_j \varphi_j$ as well as on plotting u and f for visual comparison. quantitative comparison needs to investigate the error e(x) = f(x) - u mostly want a single number to reflect the error and use a norm for this jusually the L^2 norm

$$||e||_{L^2} = \left(\int_{\Omega} e^2 dx\right)^{1/2}$$
.

Since the finite element approximation is defined for all $x \in \Omega$, and interested in how u(x) deviates from f(x) through all the elements, we ca integrate analytically or use an accurate numerical approximation. The more convenient as it is a generally feasible and simple approach. The ic sample e(x) at a large number of points in each element. The function in the fe_approx1D_numint module does this for u(x) and returns an with coordinates and an array u with the u values:

Let us use the Trapezoidal method to approximate the integral. Because elements may have different lengths, the x array has a non-uniformly disset of coordinates. Also, the u_glob function works in an element by fashion such that coordinates at the boundaries between elements appeated the trapezoidal rule where add up all the trapezoids:

$$\int_{\Omega} g(x)dx \approx \sum_{j=0}^{n-1} \frac{1}{2} (g(x_j) + g(x_{j+1}))(x_{j+1} - x_j),$$

if x_0, \ldots, x_n are all the coordinates in x. In vectorized Python code,

```
g_x = g(x)
integral = 0.5*np.sum((g_x[:-1] + g_x[1:])*(x[1:] - x[:-1]))
```

Computing the L^2 norm of the error, here named E, is now achieved by

```
e2 = e**2
E = np.sqrt(0.5*np.sum((e2[:-1] + e2[1:])*(x[1:] - x[:-1]))
```

How does the error depend on h and d?

Theory and experiments show that the least squares or projection/Gal method in combination with Pd elements of equal length h has an entire of equal length h has an enti

$$||e||_{L^2} = Ch^{d+1}, (93)$$

where C is a constant depending on f, but not on h or d.

.5 Example: Cubic Hermite polynomials

he finite elements considered so far represent u as piecewise polynomials with iscontinuous derivatives at the cell boundaries. Sometimes it is desirable to ave continuous derivatives. A primary examples is the solution of differential quations with fourth-order derivatives where standard finite element formulaons lead to a need for basis functions with continuous first-order derivatives. he most common type of such basis functions in 1D is the so-called cubic ermite polynomials. The construction of such polynomials, as explained next, ill further exemplify the concepts of a cell, vertex, degree of freedom, and dof lap.

Given a reference cell [-1,1], we seek cubic polynomials with the values of ne function and its first-order derivative at X=-1 and X=1 as the four egrees of freedom. Let us number the degrees of freedom as

- 0: value of function at X = -1
- 1: value of first derivative at X = -1
- 2: value of function at X = 1
- 3: value of first derivative at X=1

y having the derivatives as unknowns, we ensure that the derivative of a basis unction in two neighboring elements is the same at the node points.

The four basis functions can be written in a general form

$$\tilde{\varphi}_i(X) = \sum_{j=0}^3 C_{i,j} X^j,$$

ith four coefficients $C_{i,j}$, j=0,1,2,3, to be determined for each i. The instraints that basis function number i must be 1 for degree of freedom number and zero for the other three degrees of freedom, gives four equations to determine i, j for each j. In mathematical detail,

$$\begin{split} \tilde{\varphi}_0(-1) &= 1, \quad \tilde{\varphi}_0(1) = \tilde{\varphi}_0'(-1) = \tilde{\varphi}_i'(1) = 0, \\ \tilde{\varphi}_1'(-1) &= 1, \quad \tilde{\varphi}_1(-1) = \tilde{\varphi}_1(1) = \tilde{\varphi}_1'(1) = 0, \\ \tilde{\varphi}_2(1) &= 1, \quad \tilde{\varphi}_2(-1) = \tilde{\varphi}_2'(-1) = \tilde{\varphi}_2'(1) = 0, \\ \tilde{\varphi}_3'(1) &= 1, \quad \tilde{\varphi}_3(-1) = \tilde{\varphi}_3'(-1) = \tilde{\varphi}_3(1) = 0. \end{split}$$

hese four 4×4 linear equations can be solved, yielding the following formulas or the cubic basis functions:

$$\tilde{\varphi}_0(X) = 1 - \frac{3}{4}(X+1)^2 + \frac{1}{4}(X+1)^3$$

$$\tilde{\varphi}_1(X) = -(X+1)(1 - \frac{1}{2}(X+1))^2$$

$$\tilde{\varphi}_2(X) = \frac{3}{4}(X+1)^2 - \frac{1}{2}(X+1)^3$$

$$\tilde{\varphi}_3(X) = -\frac{1}{2}(X+1)(\frac{1}{2}(X+1)^2 - (X+1))$$

The construction of the dof map needs a scheme for numbering the degrees of freedom. A natural left-to-right numbering has the function vertex x_i as degree of freedom number 2i and the value of the derivati as degree of freedom number 2i + 1, $i = 0, \ldots, N_e + 1$.

7 Numerical integration

Finite element codes usually apply numerical approximations to integral the integrands in the coefficient matrix often are (lower-order) polyn integration rules that can integrate polynomials exactly are popular.

The numerical integration rules can be expressed in a common form

$$\int_{-1}^{1} g(X)dX \approx \sum_{j=0}^{M} w_j g(\bar{X}_j),$$

where \bar{X}_j are integration points and w_j are integration weights, j=0 Different rules correspond to different choices of points and weights.

The very simplest method is the *Midpoint rule*,

$$\int_{-1}^{1} g(X)dX \approx 2g(0), \quad \bar{X}_{0} = 0, \ w_{0} = 2,$$

which integrates linear functions exactly.

7.1 Newton-Cotes rules

The Newton-Cotes⁸ rules are based on a fixed uniform distribution integration points. The first two formulas in this family are the well *Trapezoidal rule*,

$$\int_{-1}^{1} g(X)dX \approx g(-1) + g(1), \quad \bar{X}_{0} = -1, \ \bar{X}_{1} = 1, \ w_{0} = w_{1} = 1,$$

⁸http://en.wikipedia.org/wiki/Newton%E2%80%93Cotes formulas

nd Simpson's rule,

$$\int_{-1}^{1} g(X)dX \approx \frac{1}{3} \left(g(-1) + 4g(0) + g(1) \right), \tag{102}$$

here

$$\bar{X}_0 = -1, \ \bar{X}_1 = 0, \ \bar{X}_2 = 1, \ w_0 = w_2 = \frac{1}{3}, \ w_1 = \frac{4}{3}.$$
 (103)

ewton-Cotes rules up to five points is supported in the module file numint.py⁹. For higher accuracy one can divide the reference cell into a set of subintervals nd use the rules above on each subinterval. This approach results in *composite* iles, well-known from basic introductions to numerical integration of $\int_a^b f(x)dx$.

.2 Gauss-Legendre rules with optimized points

lore accurate rules, for a given M, arise if the location of the integration points re optimized for polynomial integrands. The Gauss-Legendre rules 10 (also nown as Gauss-Legendre quadrature or Gaussian quadrature) constitute one ich class of integration methods. Two widely applied Gauss-Legendre rules in its family have the choice

$$M = 1: \quad \bar{X}_0 = -\frac{1}{\sqrt{3}}, \ \bar{X}_1 = \frac{1}{\sqrt{3}}, \ w_0 = w_1 = 1$$
 (104)

$$M = 2: \quad \bar{X}_0 = -\sqrt{\frac{3}{5}}, \ \bar{X}_0 = 0, \ \bar{X}_2 = \sqrt{\frac{3}{5}}, \ w_0 = w_2 = \frac{5}{9}, \ w_1 = \frac{8}{9}.$$
 (105)

hese rules integrate 3rd and 5th degree polynomials exactly. In general, an I-point Gauss-Legendre rule integrates a polynomial of degree 2M+1 exactly. he code numint.py contains a large collection of Gauss-Legendre rules.

Approximation of functions in 2D

Il the concepts and algorithms developed for approximation of 1D functions (x) can readily be extended to 2D functions f(x,y) and 3D functions f(x,y,z). asically, the extensions consists of defining basis functions $\psi_i(x,y)$ or $\psi_i(x,y,z)$ ver some domain Ω , and for the least squares and Galerkin methods, the itegration is done over Ω .

As in 1D, the least squares and projection/Galerkin methods two lead to near systems $\,$

$$\sum_{j \in \mathcal{I}_s} A_{i,j} c_j = b_i, \quad i \in \mathcal{I}_s,$$
$$A_{i,j} = (\psi_i, \psi_j),$$
$$b_i = (f, \psi_i),$$

where the inner product of two functions f(x, y) and g(x, y) is defined coranalogously to the 1D case (24):

$$(f,g) = \int_{\Omega} f(x,y)g(x,y)dxdy$$

8.1 2D basis functions as tensor products of 1D fun-

One straightforward way to construct a basis in 2D is to combine 1 functions. Say we have the 1D vector space

$$V_x = \operatorname{span}\{\hat{\psi}_0(x), \dots, \hat{\psi}_{N_x}(x)\}.$$

A similar space for variation in y can be defined,

$$V_y = \operatorname{span}\{\hat{\psi}_0(y), \dots, \hat{\psi}_{N_y}(y)\}.$$

We can then form 2D basis functions as tensor products of 1D basis fur

Tensor products.

Given two vectors $a = (a_0, \ldots, a_M)$ and $b = (b_0, \ldots, b_N)$, their outer to product, also called the dyadic product, is $p = a \otimes b$, defined through

$$p_{i,j} = a_i b_j, \quad i = 0, \dots, M, \ j = 0, \dots, N.$$

In the tensor terminology, a and b are first-order tensors (vectors with index, also termed rank-1 tensors), and then their outer tensor product a second-order tensor (matrix with two indices, also termed rank-2 tensor product is the well-known scalar of product of two vectors: $p = a \cdot b = \sum_{j=0}^{N} a_j b_j$. Now, p is a rank-0 tensor product of two vectors:

Tensors are typically represented by arrays in computer code. In above example, a and b are represented by one-dimensional array length M and N, respectively, while $p = a \otimes b$ must be represented two-dimensional array of size $M \times N$.

Tensor products^a can be used in a variety of context.

Given the vector spaces V_x and V_y as defined in (107) and (108), the product space $V = V_x \otimes V_y$ has a basis formed as the tensor product

⁹http://tinyurl.com/nm5587k/fem/numint.py

¹⁰http://en.wikipedia.org/wiki/Gaussian_quadrature

ahttp://en.wikipedia.org/wiki/Tensor_product

asis for V_x and V_y . That is, if $\{\varphi_i(x)\}_{i\in\mathcal{I}_x}$ and $\{\varphi_i(y)\}_{i\in\mathcal{I}_y}$ are basis for V_x and V_y , respectively, the elements in the basis for V arise from the tensor roduct: $\{\varphi_i(x)\varphi_j(y)\}_{i\in\mathcal{I}_x,j\in\mathcal{I}_y}$. The index sets are $I_x=\{0,\ldots,N_x\}$ and $I_y=\{0,\ldots,N_y\}$.

The notation for a basis function in 2D can employ a double index as in

$$\psi_{p,q}(x,y) = \hat{\psi}_p(x)\hat{\psi}_q(y), \quad p \in \mathcal{I}_x, q \in \mathcal{I}_y.$$

he expansion for u is then written as a double sum

$$u = \sum_{p \in \mathcal{I}_x} \sum_{q \in \mathcal{I}_y} c_{p,q} \psi_{p,q}(x,y).$$

lternatively, we may employ a single index,

$$\psi_i(x,y) = \hat{\psi}_p(x)\hat{\psi}_q(y),$$

nd use the standard form for u,

$$u = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x, y) .$$

he single index is related to the double index through $i = p(N_y + 1) + q$ or $q(N_x + 1) + p$.

.2 Example: Polynomial basis in 2D

uppose we choose $\hat{\psi}_p(x) = x^p$, and try an approximation with $N_x = N_y = 1$:

$$\psi_{0,0} = 1$$
, $\psi_{1,0} = x$, $\psi_{0,1} = y$, $\psi_{1,1} = xy$.

sing a mapping to one index like $i = q(N_x + 1) + p$, we get

$$\psi_0 = 1$$
, $\psi_1 = x$, $\psi_2 = y$, $\psi_3 = xy$.

With the specific choice $f(x,y) = (1+x^2)(1+2y^2)$ on $\Omega = [0,L_x] \times [0,L_y]$ e can perform actual calculations:

$$\begin{split} A_{0,0} &= (\psi_0, \psi_0) = \int_0^{L_y} \int_0^{L_x} \psi_0(x,y)^2 dx dy = \int_0^{L_y} \int_0^{L_x} dx dy = L_x L_y, \\ A_{1,0} &= (\psi_1, \psi_0) = \int_0^{L_y} \int_0^{L_x} x dx dy = \frac{1}{2} L_x^2 L_y, \\ A_{0,1} &= (\psi_0, \psi_1) = \int_0^{L_y} \int_0^{L_x} y dx dy = \frac{1}{2} L_y^2 L_x, \\ A_{0,1} &= (\psi_0, \psi_1) = \int_0^{L_y} \int_0^{L_x} x y dx dy = \int_0^{L_y} y dy \int_0^{L_x} x dx = \frac{1}{4} L_y^2 L_x^2. \end{split}$$

he right-hand side vector has the entries

$$b_0 = (\psi_0, f) = \int_0^{L_y} \int_0^{L_x} 1 \cdot (1 + x^2)(1 + 2y^2) dx dy$$

$$= \int_0^{L_y} (1 + 2y^2) dy \int_0^{L_x} (1 + x^2) dx = (L_y + \frac{2}{3}L_y^3)(L_x + \frac{1}{3}L_x^3)$$

$$b_1 = (\psi_1, f) = \int_0^{L_y} \int_0^{L_x} x(1 + x^2)(1 + 2y^2) dx dy$$

$$= \int_0^{L_y} (1 + 2y^2) dy \int_0^{L_x} x(1 + x^2) dx = (L_y + \frac{2}{3}L_y^3)(\frac{1}{2}L_x^2 + \frac{1}{4}L_x^4)$$

$$b_2 = (\psi_2, f) = \int_0^{L_y} \int_0^{L_x} y(1 + x^2)(1 + 2y^2) dx dy$$

$$= \int_0^{L_y} y(1 + 2y^2) dy \int_0^{L_x} (1 + x^2) dx = (\frac{1}{2}L_y + \frac{1}{2}L_y^4)(L_x + \frac{1}{3}L_x^3)$$

$$b_3 = (\psi_2, f) = \int_0^{L_y} \int_0^{L_x} xy(1 + x^2)(1 + 2y^2) dx dy$$

$$= \int_0^{L_y} y(1 + 2y^2) dy \int_0^{L_x} xy(1 + x^2) dx = (\frac{1}{2}L_y^2 + \frac{1}{2}L_y^4)(\frac{1}{2}L_x^2 + \frac{1}{4}I_x^4)$$

There is a general pattern in these calculations that we can exple arbitrary matrix entry has the formula

$$\begin{split} A_{i,j} &= (\psi_i, \psi_j) = \int_0^{L_y} \int_0^{L_x} \psi_i \psi_j dx dy \\ &= \int_0^{L_y} \int_0^{L_x} \psi_{p,q} \psi_{r,s} dx dy = \int_0^{L_y} \int_0^{L_x} \hat{\psi}_p(x) \hat{\psi}_q(y) \hat{\psi}_r(x) \hat{\psi}_s(y) dx \\ &= \int_0^{L_y} \hat{\psi}_q(y) \hat{\psi}_s(y) dy \int_0^{L_x} \hat{\psi}_p(x) \hat{\psi}_r(x) dx \\ &= \hat{A}_{p,r}^{(x)} \hat{A}_{q,s}^{(y)}, \end{split}$$

where

$$\hat{A}_{p,r}^{(x)} = \int_{0}^{L_{x}} \hat{\psi}_{p}(x) \hat{\psi}_{r}(x) dx, \quad \hat{A}_{q,s}^{(y)} = \int_{0}^{L_{y}} \hat{\psi}_{q}(y) \hat{\psi}_{s}(y) dy,$$

are matrix entries for one-dimensional approximations. Moreover, i=1 and $j=sN_y+r$.

With $\hat{\psi}_p(x) = x^p$ we have

$$\hat{A}_{p,r}^{(x)} = \frac{1}{n+r+1} L_x^{p+r+1}, \quad \hat{A}_{q,s}^{(y)} = \frac{1}{q+s+1} L_y^{q+s+1},$$

and

$$A_{i,j} = \hat{A}_{p,r}^{(x)} \hat{A}_{q,s}^{(y)} = \frac{1}{p+r+1} L_x^{p+r+1} \frac{1}{q+s+1} L_y^{q+s+1},$$

or $p, r \in \mathcal{I}_x$ and $q, s \in \mathcal{I}_y$.

Corresponding reasoning for the right-hand side leads to

$$b_{i} = (\psi_{i}, f) = \int_{0}^{L_{y}} \int_{0}^{L_{x}} \psi_{i} f \, dx dx$$

$$= \int_{0}^{L_{y}} \int_{0}^{L_{x}} \hat{\psi}_{p}(x) \hat{\psi}_{q}(y) f \, dx dx$$

$$= \int_{0}^{L_{y}} \hat{\psi}_{q}(y) (1 + 2y^{2}) dy \int_{0}^{L_{y}} \hat{\psi}_{p}(x) x^{p} (1 + x^{2}) dx$$

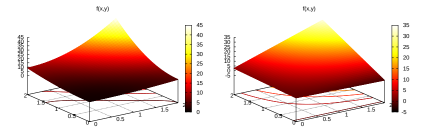
$$= \int_{0}^{L_{y}} y^{q} (1 + 2y^{2}) dy \int_{0}^{L_{y}} x^{p} (1 + x^{2}) dx$$

$$= (\frac{1}{q+1} L_{y}^{q+1} + \frac{2}{q+3} L_{y}^{q+3}) (\frac{1}{p+1} L_{x}^{p+1} + \frac{2}{q+3} L_{x}^{p+3})$$

Choosing $L_x = L_y = 2$, we have

$$A = \begin{bmatrix} 4 & 4 & 4 & 4 \\ 4 & \frac{16}{3} & 4 & \frac{16}{3} \\ 4 & 4 & \frac{16}{3} & \frac{16}{6} & \frac{64}{3} \\ 4 & \frac{16}{3} & \frac{16}{3} & \frac{64}{4} \end{bmatrix}, \quad b = \begin{bmatrix} \frac{308}{9} \\ \frac{140}{3} \\ 44 \\ 60 \end{bmatrix}, \quad c = \begin{bmatrix} -\frac{1}{9} \\ \frac{4}{3} \\ -\frac{2}{3} \\ 8 \end{bmatrix}.$$

igure 33 illustrates the result.



igure 33: Approximation of a 2D quadratic function (left) by a 2D bilinear unction (right) using the Galerkin or least squares method.

.3 Implementation

he least_squares function from Section 2.8 and/or the file approx1D.py¹¹ an with very small modifications solve 2D approximation problems. First,

let Omega now be a list of the intervals in x and y direction. For e $\Omega = [0, L_x] \times [0, L_y]$ can be represented by Omega = [[0, L_x], [0, 1 Second, the symbolic integration must be extended to 2D:

provided integrand is an expression involving the ${\tt sympy}$ symbols x and 2D version of numerical integration becomes

The right-hand side integrals are modified in a similar way.

Third, we must construct a list of 2D basis functions. Here are two expansed on tensor products of 1D "Taylor-style" polynomials x^i and functions $\sin((i+1)\pi x)$:

The complete code appears in approx2D.py¹².

The previous hand calculation where a quadratic f was approximat bilinear function can be computed symbolically by

```
>>> from approx2D import *
>>> f = (1+x*2)*(1+2*y**2)
>>> psi = taylor(x, y, 1, 1)
>>> Omega = [[0, 2], [0, 2]]
>>> u, c = least_squares(f, psi, Omega)
>>> print u
8*x*y - 2*x/3 + 4*y/3 - 1/9
>>> print sp.expand(f)
2*x**2*y**2 + x**2 + 2*y**2 + 1
```

We may continue with adding higher powers to the basis:

```
>>> psi = taylor(x, y, 2, 2)
>>> u, c = least_squares(f, psi, Omega)
>>> print u
2*x**2*y**2 + x**2 + 2*y**2 + 1
>>> print u-f
0
```

¹¹http://tinyurl.com/nm5587k/fem/fe_approx1D.py

¹²http://tinyurl.com/nm5587k/fem/fe_approx2D.py

or $N_x \ge 2$ and $N_y \ge 2$ we recover the exact function f, as expected, since in at case $f \in V$ (see Section 2.5).

.4 Extension to 3D

xtension to 3D is in principle straightforward once the 2D extension is unerstood. The only major difference is that we need the repeated outer tensor roduct.

$$V = V_x \otimes V_y \otimes V_z$$
.

ı general, given vectors (first-order tensors) $a^{(q)}=(a_0^{(q)},\ldots,a_{N_q}^{(q)},\,q=0,\ldots,m,$ ne tensor product $p=a^{(0)}\otimes\cdots\otimes a^m$ has elements

$$p_{i_0,i_1,\ldots,i_m} = a_{i_1}^{(0)} a_{i_1}^{(1)} \cdots a_{i_m}^{(m)}.$$

he basis functions in 3D are then

$$\psi_{p,q,r}(x,y,z) = \hat{\psi}_p(x)\hat{\psi}_q(y)\hat{\psi}_r(z),$$

ith $p \in \mathcal{I}_x$, $q \in \mathcal{I}_y$, $r \in \mathcal{I}_z$. The expansion of u becomes

$$u(x, y, z) = \sum_{p \in \mathcal{I}_x} \sum_{q \in \mathcal{I}_y} \sum_{r \in \mathcal{I}_z} c_{p,q,r} \psi_{p,q,r}(x, y, z).$$

single index can be introduced also here, e.g., $i=N_xN_yr+q_Nx+p,\ u=\sum_i c_i\psi_i(x,y,z).$

Use of tensor product spaces.

Constructing a multi-dimensional space and basis from tensor products of 1D spaces is a standard technique when working with global basis functions. In the world of finite elements, constructing basis functions by tensor products is much used on quadrilateral and hexahedra cell shapes, but not on triangles and tetrahedra. Also, the global finite element basis functions are almost exclusively denoted by a single index and not by the natural tuple of indices that arises from tensor products.

Finite elements in 2D and 3D

inite element approximation is particularly powerful in 2D and 3D because ne method can handle a geometrically complex domain Ω with ease. The rincipal idea is, as in 1D, to divide the domain into cells and use polynomials or approximating a function over a cell. Two popular cell shapes are triangles nd the quadrilaterals. Figures 34, 35, and 36 provide examples. P1 elements

means linear functions $(a_0 + a_1x + a_2y)$ over triangles, while Q1 elements bilinear functions $(a_0 + a_1x + a_2y + a_3xy)$ over rectangular cells. Higher elements can easily be defined.

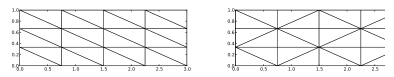


Figure 34: Examples on 2D P1 elements.

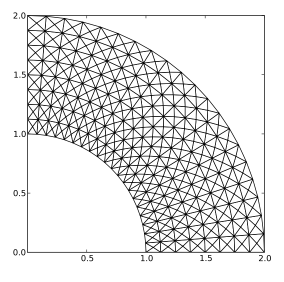


Figure 35: Examples on 2D P1 elements in a deformed geometry

9.1 Basis functions over triangles in the physical do

Cells with triangular shape will be in main focus here. With the P1 tri element, u is a linear function over each cell, as depicted in Figure 3 discontinuous derivatives at the cell boundaries.

We give the vertices of the cells global and local numbers as in 1 degrees of freedom in the P1 element are the function values at a set of which are the three vertices. The basis function $\varphi_i(x,y)$ is then 1 at the with global vertex number i and zero at all other vertices. On an element three degrees of freedom uniquely determine the linear basis functions

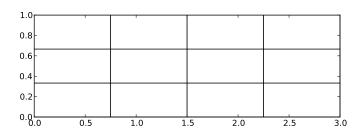


Figure 36: Examples on 2D Q1 elements.

ement, as usual. The global $\varphi_i(x,y)$ function is then a combination of the linear nections (planar surfaces) over all the neighboring cells that have vertex number in common. Figure 38 tries to illustrate the shape of such a "pyramid"-like nection.

Element matrices and vectors. As in 1D, we split the integral over Ω into sum of integrals over cells. Also as in 1D, φ_i overlaps φ_j (i.e., $\varphi_i\varphi_j\neq 0$) if nd only if i and j are vertices in the same cell. Therefore, the integral of $\varphi_i\varphi_j$ ver an element is nonzero only when i and j run over the vertex numbers in ne element. These nonzero contributions to the coefficient matrix are, as in 1D, ollected in an element matrix. The size of the element matrix becomes 3×3 nce there are three degrees of freedom that i and j run over. Again, as in 1D, e number the local vertices in a cell, starting at 0, and add the entries in the ement matrix into the global system matrix, exactly as in 1D. All details and ode appear below.

.2 Basis functions over triangles in the reference cell

s in 1D, we can define the basis functions and the degrees of freedom in a eference cell and then use a mapping from the reference coordinate system to rephysical coordinate system. We also have a mapping of local degrees of eedom numbers to global degrees of freedom numbers.

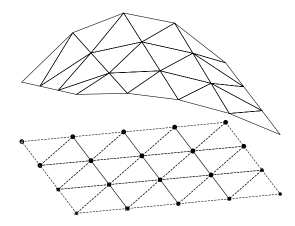


Figure 37: Example on piecewise linear 2D functions defined on trial

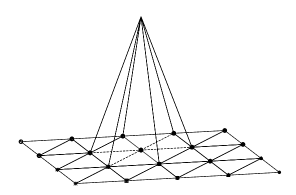
The reference cell in an (X,Y) coordinate system has vertices (0,0) and (0,1), corresponding to local vertex numbers (0,1), and (0,1), respective P1 element has linear functions $\tilde{\varphi}_r(X,Y)$ as basis functions, (r) = (0,1,2). linear function $\tilde{\varphi}_r(X,Y)$ in 2D is on the form $(C_{r,0} + C_{r,1}X + C_{r,2}Y)$, and has three parameters $(C_{r,0}, C_{r,1})$, and $(C_{r,2})$, we need three degrees of f These are in general taken as the function values at a set of nodes. For element the set of nodes is the three vertices. Figure 39 displays the go of the element and the location of the nodes.

Requiring $\tilde{\varphi}_r = 1$ at node number r and $\tilde{\varphi}_r = 0$ at the two other node three linear equations to determine $C_{r,0}$, $C_{r,1}$, and $C_{r,2}$. The result is

$$\begin{split} \tilde{\varphi}_0(X,Y) &= 1 - X - Y, \\ \tilde{\varphi}_1(X,Y) &= X, \\ \tilde{\varphi}_2(X,Y) &= Y \end{split}$$

Higher-order approximations are obtained by increasing the polynomi adding additional nodes, and letting the degrees of freedom be function at the nodes. Figure 40 shows the location of the six nodes in the P2 e

A polynomial of degree p in X and Y has $n_p = (p+1)(p+2)/2$ tendence needs n_p nodes. The values at the nodes constitute n_p degrees of $\tilde{\varphi}_r$ up to degree 6 is displayed in Figure 4



igure 38: Example on a piecewise linear 2D basis function over a patch of riangles.

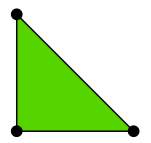


Figure 39: 2D P1 element.

The generalization to 3D is straightforward: the reference element is a strahedron 13 with vertices $(0,0,0),\,(1,0,0),\,(0,1,0),\,$ and (0,0,1) in a X,Y,Z eference coordinate system. The P1 element has its degrees of freedom as four odes, which are the four vertices, see Figure 42. The P2 element adds additional odes along the edges of the cell, yielding a total of 10 nodes and degrees of eedom, see Figure 43.

The interval in 1D, the triangle in 2D, the tetrahedron in 3D, and its eneralizations to higher space dimensions are known as *simplex* cells (the eometry) or *simplex* elements (the geometry, basis functions, degrees of freedom,

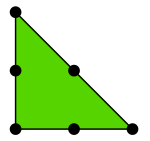


Figure 40: 2D P2 element.

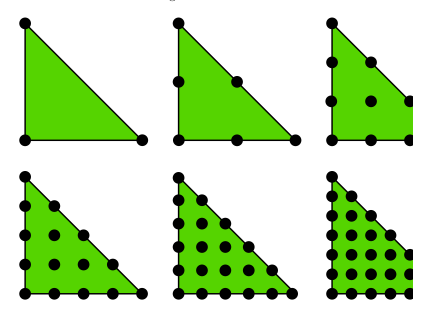


Figure 41: 2D P1, P2, P3, P4, P5, and P6 elements.

etc.). The plural forms simplices 14 and simplexes are also a much used terms when referring to this type of cells or elements. The side of a sir called a *face*, while the tetrahedron also has *edges*.

Acknowledgment. Figures 39 to 43 are created by Anders Logg an from the FEniCS book¹⁵: *Automated Solution of Differential Equation Finite Element Method*, edited by A. Logg, K.-A. Mardal, and G. N published by Springer¹⁶, 2012.

¹³http://en.wikipedia.org/wiki/Tetrahedron

¹⁴http://en.wikipedia.org/wiki/Simplex

¹⁵https://launchpad.net/fenics-book

¹⁶http://goo.gl/lbyVMH

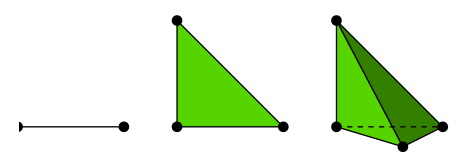


Figure 42: P1 elements in 1D, 2D, and 3D.

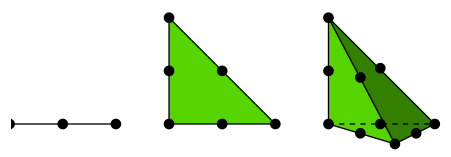


Figure 43: P2 elements in 1D, 2D, and 3D.

.3 Affine mapping of the reference cell

et $\tilde{\varphi}_r^{(1)}$ denote the basis functions associated with the P1 element in 1D, 2D, or D, and let $\boldsymbol{x}_{q(e,r)}$ be the physical coordinates of local vertex number r in cell e. urthermore, let \boldsymbol{X} be a point in the reference coordinate system corresponding the point \boldsymbol{x} in the physical coordinate system. The affine mapping of any \boldsymbol{X} nto \boldsymbol{x} is then defined by

$$\boldsymbol{x} = \sum_{r} \tilde{\varphi}_{r}^{(1)}(\boldsymbol{X}) \boldsymbol{x}_{q(e,r)}, \tag{112}$$

here r runs over the local vertex numbers in the cell. The affine mapping sentially stretches, translates, and rotates the triangle. Straight or planar ices of the reference cell are therefore mapped onto straight or planar faces in the physical coordinate system. The mapping can be used for both P1 and igher-order elements, but note that the mapping itself always applies the P1 axis functions.

.4 Isoparametric mapping of the reference cell

istead of using the P1 basis functions in the mapping (112), we may use the asis functions of the actual Pd element:

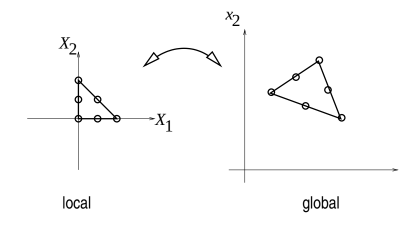


Figure 44: Affine mapping of a P1 element.

$$oldsymbol{x} = \sum_r ilde{arphi}_r(oldsymbol{X}) oldsymbol{x}_{q(e,r)},$$

where r runs over all nodes, i.e., all points associated with the degrees of This is called an *isoparametric mapping*. For P1 elements it is iden the affine mapping (112), but for higher-order elements the mapping straight or planar faces of the reference cell will result in a *curved* face physical coordinate system. For example, when we use the basis function triangular P2 element in 2D in (113), the straight faces of the reference are mapped onto curved faces of parabolic shape in the physical coc system, see Figure 45.

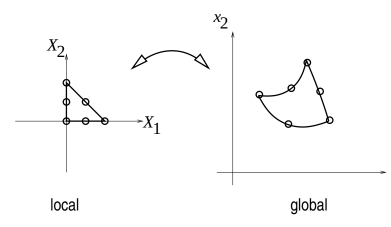


Figure 45: Isoparametric mapping of a P2 element.

From (112) or (113) it is easy to realize that the vertices are correctly mapped. onsider a vertex with local number s. Then $\tilde{\varphi}_s = 1$ at this vertex and zero at 12 others. This means that only one term in the sum is nonzero and $\boldsymbol{x} = \boldsymbol{x}_{q(e,s)}$, hich is the coordinate of this vertex in the global coordinate system.

.5 Computing integrals

et $\tilde{\Omega}^r$ denote the reference cell and $\Omega^{(e)}$ the cell in the physical coordinate vstem. The transformation of the integral from the physical to the reference pordinate system reads

$$\int_{\Omega^{(e)}} \varphi_i(\boldsymbol{x}) \varphi_j(\boldsymbol{x}) \, \mathrm{d}\boldsymbol{x} = \int_{\tilde{\Omega}^r} \tilde{\varphi}_i(\boldsymbol{X}) \tilde{\varphi}_j(\boldsymbol{X}) \, \mathrm{det} \, J \, \, \mathrm{d}\boldsymbol{X}, \tag{114}$$

$$\int_{\Omega^{(e)}} \varphi_i(\boldsymbol{x}) f(\boldsymbol{x}) \, \mathrm{d}\boldsymbol{x} = \int_{\tilde{\Omega}^r} \tilde{\varphi}_i(\boldsymbol{X}) f(\boldsymbol{x}(\boldsymbol{X})) \, \mathrm{det} \, J \, \, \mathrm{d}\boldsymbol{X}, \tag{115}$$

here dx means the infinitesimal area element dxdy in 2D and dxdydz in 3D, ith a similar definition of dX. The quantity $\det J$ is the determinant of the acobian of the mapping $\boldsymbol{x}(X)$. In 2D,

$$J = \begin{bmatrix} \frac{\partial x}{\partial X} & \frac{\partial x}{\partial Y} \\ \frac{\partial y}{\partial X} & \frac{\partial y}{\partial Y} \end{bmatrix}, \quad \det J = \frac{\partial x}{\partial X} \frac{\partial y}{\partial Y} - \frac{\partial x}{\partial Y} \frac{\partial y}{\partial X}.$$
 (116)

/ith the affine mapping (112), $\det J = 2\Delta$, where Δ is the area or volume of ne cell in the physical coordinate system.

temark. Observe that finite elements in 2D and 3D builds on the same *ideas* and *concepts* as in 1D, but there is simply much more to compute because the pecific mathematical formulas in 2D and 3D are more complicated and the book eeping with dof maps also gets more complicated. The manual work is tedious, engthy, and error-prone so automation by the computer is a must.

0 Exercises

xercise 1: Linear algebra refresher I

ook up the topic of *vector space* in your favorite linear algebra book or search or the term at Wikipedia. Prove that vectors in the plane (a,b) form a vector pace by showing that all the axioms of a vector space are satisfied. Similarly, rove that all linear functions of the form ax + b constitute a vector space, $b \in \mathbb{R}$.

On the contrary, show that all quadratic functions of the form $1 + ax^2 + bx$ on not constitute a vector space. Filename: linalg1.pdf.

Exercise 2: Linear algebra refresher II

As an extension of Exercise 1, check out the topic of *inner product spaces*. a possible inner product for the space of all linear functions of the form $a,b \in \mathbb{R}$. Show that this inner product satisfies the general requiremen inner product in a vector space. Filename: linalg2.pdf.

Exercise 3: Approximate a three-dimensional vector plane

Given $\mathbf{f} = (1,1,1)$ in \mathbb{R}^3 , find the best approximation vector \mathbf{u} in the spanned by the unit vectors (1,0) and (0,1). Repeat the calculations unvectors (2,1) and (1,2). Filename: vec111_approx.pdf.

Exercise 4: Approximate the exponential function by j functions

Let V be a function space with basis functions x^i , i = 0, 1, ..., N. Find approximation to $f(x) = \exp(-x)$ on $\Omega = [0, 8]$ among all functions i N = 2, 4, 6. Illustrate the three approximations in three separate plots.

Hint. The exercise is easy to solve if you apply the <code>lest_squares</code> and <code>cc</code> functions in the <code>approx1D.py</code> module.

Filename: exp_powers.py.

Exercise 5: Approximate the sine function by power tions

In this exercise we want to approximate the sine function by polynor order N+1. Consider two bases:

$$V_1 = \{x, x^3, x^5, \dots, x^{N-2}, x^N\},$$

$$V_2 = \{1, x, x^2, x^3, \dots, x^N\}.$$

The basis V_1 is motivated by the fact that the Taylor polynomial approx to the sine function has only odd powers, while V_2 is motivated by the assithat also the even powers could improve the approximation in a least-setting.

Compute the best approximation to $f(x) = \sin(x)$ among all func V_1 and V_2 on two domains of increasing sizes: $\Omega_{1,k} = [0,k\pi], \ k=2,3\dots$ $\Omega_{2,k} = [-k\pi/2,k\pi/2], \ k=2,3\dots,6$. Make plots for all combinations o $\Omega_1, \Omega_2, \ k=2,3,\dots,6$.

Add a plot of the N-th degree Taylor polynomial approximation α around x = 0.

lint. You can make a loop over V_1 and V_2 , a loop over Ω_1 and Ω_2 , and a loop ver k. Inside the loops, call the functions least_squares and comparison_plot om the approx1D module. N=9 is a suggested value. ilename: \sin_p owers.py.

Exercise 6: Approximate a steep function by sines

ind the best approximation of $f(x) = \tanh(s(x-\pi))$ on $[0,2\pi]$ in the space V ith basis $\psi_i(x) = \sin((2i+1)x)$, $i \in \mathcal{I}_s = \{0,\ldots,N\}$. Make a movie showing ow $u = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x)$ approximates f(x) as N grows. Choose s such that f is seep (s = 20) may be appropriate).

lint. One may naively call the least_squares_orth and comparison_plot om the approx1D module in a loop and extend the basis with one new element 1 each pass. This approach implies a lot of recomputations. A more efficient rategy is to let least_squares_orth compute with only one basis function at time and accumulate the corresponding u in the total solution. ilename: tanh_sines_approx.py.

temarks. Approximation of a discontinuous (or steep) f(x) by sines, results a slow convergence and oscillatory behavior of the approximation close to the brupt changes in f. This is known as the Gibb's phenomenon¹⁷.

Exercise 7: Animate the approximation of a steep function by sines

lake a movie that shows how the approximation in Exercise 6 is improved as 7 grows. Illustrate a smooth case where s=0.5 and a steep case where s=20 the $\tanh(s(x-\pi))$ function. Filename: $\tanh_sines_approx_movie.py$.

'exercise 8: Fourier series as a least squares approximation

iven a function f(x) on an interval [0, L], look up the formula for the coefficients i and b_i in the Fourier series of f:

$$f(x) = a_0 + \sum_{j=1}^{\infty} a_j \cos\left(j\frac{\pi x}{L}\right) + \sum_{j=1}^{\infty} b_j \sin\left(j\frac{\pi x}{L}\right).$$

Let an infinite-dimensional vector space V have the basis functions $\cos j \frac{\pi x}{L}$ or $j=0,1,\ldots,\infty$ and $\sin j \frac{\pi x}{L}$ for $j=1,\ldots,\infty$. Show that the least squares pproximation method from Section 2 leads to a linear system whose solution pincides with the standard formulas for the coefficients in a Fourier series of (x) (see also Section 2.7). You may choose

 $\psi_{2i} = \cos\left(i\frac{\pi}{L}x\right), \quad \psi_{2i+1} = \sin\left(i\frac{\pi}{L}x\right),$

for $i = 0, 1, \dots, N \to \infty$.

Choose $f(x) = \tanh(s(x-\frac{1}{2}))$ on $\Omega = [0,1]$, which is a smooth funct with considerable steepness around x = 1/2 as s grows in size. Calcu coefficients in the Fourier expansion by solving the linear system, arisin the least squares or Galerkin methods, by hand. Plot some truncated of the series together with f(x) to show how the series expansion conversed and s = 10 and s = 100. Filename: Fourier approx.py.

Exercise 9: Approximate a steep function by Lagrange nomials

Use interpolation/collocation with uniformly distributed points and Ch nodes to approximate

$$f(x) = -\tanh(s(x - \frac{1}{2})), \quad x \in [0, 1],$$

by Lagrange polynomials for s=10 and s=100, and N=3,6,9,11 separate plots of the approximation for each combination of s, poi (Chebyshev or uniform), and N. Filename: tanh_Lagrange.py.

Exercise 10: Define nodes and elements

Consider a domain $\Omega = [0, 2]$ divided into the three P2 elements [0, 1], and [1.2, 2].

For P1 and P2 elements, set up the list of coordinates and nodes and the numbers of the nodes that belong to each element (elements cases: 1) nodes and elements numbered from left to right, and 2) no elements numbered from right to left. Filename: fe_numberings1.py.

Exercise 11: Define vertices, cells, and dof maps

Repeat Exercise 10, but define the data structures vertices, cells, and c instead of nodes and elements. Filename: fe_numberings2.py.

Exercise 12: Construct matrix sparsity patterns

Exercise 10 describes a element mesh with a total of five elements, but v different element and node orderings. For each of the two orderings, 5×5 matrix and fill in the entries that will be nonzero.

Hint. A matrix entry (i, j) is nonzero if i and j are nodes in the same Filename: fe_sparsity_pattern.pdf.

 $^{^{17} \}verb|http://en.wikipedia.org/wiki/Gibbs_phenomenon|$

exercise 13: Perform symbolic finite element computations

erform symbolic calculations to find formulas for the coefficient matrix and ght-hand side when approximating $f(x) = \sin(x)$ on $\Omega = [0, \pi]$ by two P1 ements of size $\pi/2$. Solve the system and compare $u(\pi/2)$ with the exact value Filename: \sin_{π}

Exercise 14: Approximate a steep function by P1 and P2 lements

liven

$$f(x) = \tanh(s(x - \frac{1}{2}))$$

se the Galerkin or least squares method with finite elements to find an pproximate function u(x). Choose s=40 and try $N_e=4,8,16$ P1 elements and $N_e=2,4,8$ P2 elements. Integrate $f\varphi_i$ numerically. Filename: anh_fe_P1P2_approx.py.

Exercise 15: Approximate a steep function by P3 and P4 lements

olve Exercise 14 using $N_e=1,2,4$ P3 and P4 elements. How will a colloca-on/interpolation method work in this case with the same number of nodes? ilename: tanh_fe_P3P4_approx.py.

exercise 16: Investigate the approximation error in finite lements

he theory (93) from Section 6.4 predicts that the error in the Pd approximation f a function should behave as h^{d+1} , where h is the length of the element. Use speriments to verify this asymptotic behavior (i.e., for small enough h). Choose tree examples: $f(x) = Ae^{-\omega x}$ on $[0, 3/\omega]$, $f(x) = A\sin(\omega x)$ on $\Omega = [0, 2\pi/\omega]$ or constant A and ω , and $f(x) = \sqrt{x}$ on [0, 1].

lint. Run a series of experiments: (h_i, E_i) , i = 0, ..., m, where E_i is the L^2 orm of the error corresponding to element length h_i . Assume an error model $i = Ch^r$ and compute r from two successive experiments:

$$r_i = \ln(E_{i+1}/E_i)/\ln(h_{i+1}/h_i), \quad i = 0, \dots, m-1.$$

lopefully, the sequence r_0, \ldots, r_{m-1} converges to the true r, and r_{m-1} can be aken as an approximation to r. Run such experiments for different d for the ifferent f(x) functions.

ilename: Pd_approx_error.py.

Exercise 17: Approximate a step function by finite election

Approximate the step function

$$f(x) = \begin{cases} 1 & 0 \le x < 1/2, \\ 2 & 1/2 \le x \ge 1/2 \end{cases}$$

by 2, 4, and 8 P1 and P2 elements. Compare approximations visually.

Hint. This f can also be expressed in terms of the Heaviside functio f(x) = H(x - 1/2). Therefore, f can be defined by

f = sp.Heaviside(x - sp.Rational(1,2))

making the approximate function in the fe_approx1D.py module an candidate to solve the problem. However, sympy does not handle sintegration with this particular integrand, and the approximate function problem when converting f to a Python function (for plotting) since He is not an available function in numpy. It is better to make special-purpor for this case or perform all calculations by hand.

Filename: Heaviside_approx_P1P2.py.

Exercise 18: 2D approximation with orthogonal func

Assume we have basis functions $\varphi_i(x,y)$ in 2D that are orthogonal su $(\varphi_i,\varphi_j)=0$ when $i\neq j$. The function least_squares in the file approx will then spend much time on computing off-diagonal terms in the compatrix that we know are zero. To speed up the computations, make a least_squares_orth that utilizes the orthogonality among the basis for Apply the function to approximate

$$f(x,y) = x(1-x)y(1-y)e^{-x-y}$$

on $\Omega = [0,1] \times [0,1]$ via basis functions

$$\varphi_i(x,y) = \sin((p+1)\pi x)\sin((q+1)\pi y), \quad i = q(N_x+1) + p,$$

where $p = 0, \ldots, N_x$ and $q = 0, \ldots, N_y$.

Hint. Get ideas from the function least_squares_orth in Section file approx1D.py¹⁹.

Filename: approx2D least squares orth.py.

¹⁸http://tinyurl.com/nm5587k/fem/fe_approx2D.py

¹⁹http://tinyurl.com/nm5587k/fem/fe_approx1D.py

exercise 19: Use the Trapezoidal rule and P1 elements

onsider approximation of some f(x) on an interval Ω using the least squares or elerkin methods with P1 elements. Derive the element matrix and vector using ne Trapezoidal rule (101) for calculating integrals on the reference element. ssemble the contributions, assuming a uniform cell partitioning, and show nat the resulting linear system has the form $c_i = f(x_i)$ for $i \in \mathcal{I}_s$. Filename: e_P1_trapez.pdf.

'roblem 20: Compare P1 elements and interpolation

le shall approximate the function

$$f(x) = 1 + \epsilon \sin(2\pi nx), \quad x \in \Omega = [0, 1],$$

here $n \in \mathbb{Z}$ and $\epsilon > 0$.

-) Plot f(x) for n = 1, 2, 3 and find the wave length of the function.
-) We want to use N_P elements per wave length. Show that the number of ements is then nN_P .
-) The critical quantity for accuracy is the number of elements per wave length, of the element size in itself. It therefore suffices to study an f with just one are length in $\Omega = [0, 1]$. Set $\epsilon = 0.5$.

Run the least squares or projection/Galerkin method for $N_P=2,4,8,16,32$. Tompute the error $E=||u-f||_{L^2}$.

lint. Use the $fe_approx1D_numint$ module to compute u and use the techique from Section 6.4 to compute the norm of the error.

) Repeat the set of experiments in the above point, but use interpolation/colcation based on the node points to compute u(x) (recall that c_i is now simply (x_i)). Compute the error $E = ||u - f||_{L^2}$. Which method seems to be most courate?

ilename: P1_vs_interp.py.

exercise 21: Implement 3D computations with global basis unctions

xtend the approx2D.py²⁰ code to 3D applying ideas from Section 8.4. Use a D generalization of the test problem in Section 8.3 to test the implementation ilename: approx3D.py.

lxercise 22: Use Simpson's rule and P2 elements

edo Exercise 19, but use P2 elements and Simpson's rule based on sampling the itegrands at the nodes in the reference cell. Filename: fe_P2_simpson.pdf.

11 Basic principles for approximating diff tial equations

The finite element method is a very flexible approach for solving partial diffequations. Its two most attractive features are the ease of handling dor complex shape in two and three dimensions and the ease of constructing order discretization methods. The finite element method is usually appreciation in space, and therefore spatial problems will be our focu coming sections. Extensions to time-dependent problems may, for instantinite difference approximations in time.

Before studying how finite element methods are used to tackle diff equation, we first look at how global basis functions and the least Galerkin, and collocation principles can be used to solve differential equ

11.1 Differential equation models

Let us consider an abstract differential equation for a function u(x) variable, written as

$$\mathcal{L}(u) = 0, \quad x \in \Omega.$$

Here are a few examples on possible choices of $\mathcal{L}(u)$, of increasing comp

$$\mathcal{L}(u) = \frac{d^2 u}{dx^2} - f(x),$$

$$\mathcal{L}(u) = \frac{d}{dx} \left(\alpha(x) \frac{du}{dx} \right) + f(x),$$

$$\mathcal{L}(u) = \frac{d}{dx} \left(\alpha(u) \frac{du}{dx} \right) - au + f(x),$$

$$\mathcal{L}(u) = \frac{d}{dx} \left(\alpha(u) \frac{du}{dx} \right) + f(u, x).$$

Both $\alpha(x)$ and f(x) are considered as specified functions, while a is a preparameter. Differential equations corresponding to (118)-(119) arise in a phenomena, such as steady transport of heat in solids and flow of viscount between flat plates. The form (120) arises when transient diffusion a phenomenon are discretized in time by finite differences. The equation appear in chemical models when diffusion of a substance is combined chemical reactions. Also in biology, (121) plays an important role, it is spreading of species and in models involving generation and propagate electrical signals.

Let $\Omega = [0, L]$ be the domain in one space dimension. In addition differential equation, u must fulfill boundary conditions at the boundarie domain, x = 0 and x = L. When \mathcal{L} contains up to second-order derivations

²⁰http://tinyurl.com/nm5587k/fem/approx2D.py

the examples above, m=1, we need one boundary condition at each of the wo) boundary points, here abstractly specified as

$$\mathcal{B}_0(u) = 0, \ x = 0, \quad \mathcal{B}_1(u) = 0, \ x = L$$
 (122)

There are three common choices of boundary conditions:

$$\mathcal{B}_i(u) = u - g,$$
 Dirichlet condition (123)

$$\mathcal{B}_i(u) = -\alpha \frac{du}{dx} - g,$$
 Neumann condition (124)

$$\mathcal{B}_i(u) = -\alpha \frac{du}{dx} - h(u - g),$$
 Robin condition (125)

ere, g and a are specified quantities.

From now on we shall use $u_e(x)$ as symbol for the exact solution, fulfilling

$$\mathcal{L}(u_{\rm e}) = 0, \quad x \in \Omega, \tag{126}$$

hile u(x) is our notation for an approximate solution of the differential equation.

Remark on notation.

In the literature about the finite element method, is common to use u as the exact solution and u_h as the approximate solution, where h is a discretization parameter. However, the vast part of the present text is about the approximate solutions, and having a subscript h attached all the time is cumbersome. Of equal importance is the close correspondence between implementation and mathematics that we strive to achieve in this text: when it is natural to use ${\bf u}$ and not ${\bf u}_h$ in code, we let the mathematical notation be dictated by the code's preferred notation. After all, it is the powerful computer implementations of the finite element method that justifies studying the mathematical formulation and aspects of the method.

1.2 Simple model problems

common model problem used much in the forthcoming examples is

$$-u''(x) = f(x), \quad x \in \Omega = [0, L], \quad u(0) = 0, \ u(L) = D. \tag{127}$$

closely related problem with a different boundary condition at x=0 reads

$$-u''(x) = f(x), \quad x \in \Omega = [0, L], \quad u'(0) = C, \ u(L) = D.$$
 (128)

third variant has a variable coefficient.

$$-(\alpha(x)u'(x))' = f(x), \quad x \in \Omega = [0, L], \quad u'(0) = C, \ u(L) = D.$$

We can easily solve these using sympy. For (127) we can write the f

```
def model1(f, L, D):
    """Solve -u'' = f(x), u(0)=0, u(L)=D."""
    u_x = - sp.integrate(f, (x, 0, x)) + c_0
    u = sp.integrate(u_x, (x, 0, x)) + c_1
    r = sp.solve([u.subs(x, 0)-0, u.subs(x,L)-D], [c_0, c_1])
    u = u.subs(c_0, r[c_0]).subs(c_1, r[c_1])
    u = sp.simplify(sp.expand(u))
    return u
```

Calling model1(2, L, D) results in the solution

$$u(x) = \frac{1}{L}x\left(D + L^2 - Lx\right)$$

Model (128) can be solved by

```
def model2(f, L, C, D):
    """Solve -u'' = f(x), u'(0)=C, u(L)=D."""
    u_x = - sp.integrate(f, (x, 0, x)) + c_0
    u = sp.integrate(u_x, (x, 0, x)) + c_1
    r = sp.solve([sp.diff(u,x).subs(x, 0)-C, u.subs(x,L)-D], [c_0
    u = u.subs(c_0, r[c_0]).subs(c_1, r[c_1])
    u = sp.simplify(sp.expand(u))
    return u
```

to vield

$$u(x) = -x^2 + Cx - CL + D + L^2,$$

if f(x) = 2. Model (129) requires a bit more involved code,

```
def model3(f, a, L, C, D):
    """Solve -(a*u')' = f(x), u(0)=C, u(L)=D."""
    au_x = - sp.integrate(f, (x, 0, x)) + c_0
    u = sp.integrate(au_x/a, (x, 0, x)) + c_1
    r = sp.solve([u.subs(x, 0)-C, u.subs(x,L)-D], [c_0, c_1])
    u = u.subs(c_0, r[c_0]).subs(c_1, r[c_1])
    u = sp.simplify(sp.expand(u))
    return u
```

With f(x) = 0 and $\alpha(x) = 1 + x^2$ we get

$$u(x) = \frac{C \operatorname{atan}(L) - C \operatorname{atan}(x) + D \operatorname{atan}(x)}{\operatorname{atan}(L)}$$

1.3 Forming the residual

he fundamental idea is to seek an approximate solution u in some space V,

$$V = \operatorname{span}\{\psi_0(x), \dots, \psi_N(x)\},\$$

hich means that u can always be expressed as a linear combination of the basis inctions $\{\varphi_i\}_{i\in\mathcal{I}_e}$, with \mathcal{I}_s as the index set $\{0,\ldots,N\}$:

$$u(x) = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x) .$$

he coefficients $\{c_i\}_{i\in\mathcal{I}_s}$ are unknowns to be computed.

(Later, in Section 14, we will see that if we specify boundary values of u ifferent from zero, we must look for an approximate solution $u(x) = B(x) + \sum_j c_j \psi_j(x)$, where $\sum_j c_j \psi_j \in V$ and B(x) is some function for incorporating the right boundary values. Because of B(x), u will not necessarily lie in V. This indiffication does not imply any difficulties.)

We need principles for deriving N+1 equations to determine the N+1 nknowns $\{c_i\}_{i\in\mathcal{I}_s}$. When approximating a given function f by $u=\sum_j c_j\varphi_j$, a sy idea is to minimize the square norm of the approximation error e=u-f or quivalently) demand that e is orthogonal to V. Working with e is not so useful ere since the approximation error in our case is $e=u_e-u$ and u_e is unknown. he only general indicator we have on the quality of the approximate solution is σ what degree σ fulfills the differential equation. Inserting σ into σ in the constant σ is only likely to equal σ . The onzero result,

$$R = \mathcal{L}(u) = \mathcal{L}(\sum_{j} c_{j} \psi_{j}), \tag{132}$$

called the *residual* and measures the error in fulfilling the governing equation. Various principles for determining $\{c_i\}_{i\in\mathcal{I}_s}$ try to minimize R in some sense. ote that R varies with x and the $\{c_i\}_{i\in\mathcal{I}_s}$ parameters. We may write this ependence explicitly as

$$R = R(x; c_0, \dots, c_N)$$
 (133)

elow, we present three principles for making R small: a least squares method, projection or Galerkin method, and a collocation or interpolation method.

1.4 The least squares method

he least-squares method aims to find $\{c_i\}_{i\in\mathcal{I}_s}$ such that the square norm of the sidual

$$||R|| = (R, R) = \int_{\Omega} R^2 dx$$
 (134)

minimized. By introducing an inner product of two functions f and g on Ω as

$$(f,g) = \int_{\Omega} f(x)g(x) dx,$$

the least-squares method can be defined as

$$\min_{c_0,\dots,c_N} E = (R,R).$$

Differentiating with respect to the free parameters $\{c_i\}_{i\in\mathcal{I}_s}$ gives the equations

$$\int_{\Omega} 2R \frac{\partial R}{\partial c_i} dx = 0 \quad \Leftrightarrow \quad (R, \frac{\partial R}{\partial c_i}) = 0, \quad i \in \mathcal{I}_s.$$

11.5 The Galerkin method

The least-squares principle is equivalent to demanding the error to be ort to the space V when approximating a function f by $u \in V$. With a different equation we do not know the true error so we must instead require the R to be orthogonal to V. This idea implies seeking $\{c_i\}_{i\in\mathcal{I}_e}$ such that

$$(R, v) = 0, \quad \forall v \in V.$$

This is the Galerkin method for differential equations.

This statement is equivalent to R being orthogonal to the N+1 basis fit only:

$$(R, \psi_i) = 0, \quad i \in \mathcal{I}_s,$$

resulting in N+1 equations for determining $\{c_i\}_{i\in\mathcal{I}_s}$.

11.6 The Method of Weighted Residuals

A generalization of the Galerkin method is to demand that R is ort to some space W, but not necessarily the same space as V where we sunknown function. This generalization is naturally called the *method of residuals*:

$$(R, v) = 0, \quad \forall v \in W.$$

If $\{w_0, \ldots, w_N\}$ is a basis for W, we can equivalently express the me weighted residuals as

$$(R, w_i) = 0, \quad i \in \mathcal{I}_s.$$

The result is N+1 equations for $\{c_i\}_{i\in\mathcal{I}_s}$.

The least-squares method can also be viewed as a weighted residual with $w_i = \partial R/\partial c_i$.

Variational formulation of the continuous problem.

Formulations like (140) (or (141)) and (138) (or (139)) are known as variational formulations. These equations are in this text primarily used for a numerical approximation $u \in V$, where V is a finite-dimensional space with dimension N+1. However, we may also let V be an infinite-dimensional space containing the exact solution $u_e(x)$ such that also u_e fulfills the same variational formulation. The variational formulation is in that case a mathematical way of stating the problem and acts as an alternative to the usual formulation of a differential equation with initial and/or boundary conditions.

1.7 Test and Trial Functions

1 the context of the Galerkin method and the method of weighted residuals it is 5 mmon to use the name trial function for the approximate $u = \sum_j c_j \psi_j$. The 5 pace containing the trial function is known as the trial space. The function v attering the orthogonality requirement in the Galerkin method and the method f weighted residuals is called test function, and so are the ψ_i or w_i functions 1 at are used as weights in the inner products with the residual. The space here the test functions comes from is naturally called the test space.

We see that in the method of weighted residuals the test and trial spaces are ifferent and so are the test and trial functions. In the Galerkin method the test and trial spaces are the same (so far).

Remark.

It may be subject to debate whether it is only the form of (140) or (138) after integration by parts, as explained in Section 11.10, that qualifies for the term variational formulation. The result after integration by parts is what is obtained after taking the *first variation* of an optimization problem, see Section 11.13. However, here we use variational formulation as a common term for formulations which, in contrast to the differential equation R = 0, instead demand that an average of R is zero: (R, v) = 0 for all v in some space.

1.8 The collocation method

he idea of the collocation method is to demand that R vanishes at N+1 elected points x_0, \ldots, x_N in Ω :

$$R(x_i; c_0, \dots, c_N) = 0, \quad i \in \mathcal{I}_s.$$
(142)

The collocation method can also be viewed as a method of weighted rewith Dirac delta functions as weighting functions. Let $\delta(x-x_i)$ be the delta function centered around $x=x_i$ with the properties that $\delta(x-x_i)$ $x \neq x_i$ and

$$\int_{\Omega} f(x)\delta(x-x_i) dx = f(x_i), \quad x_i \in \Omega.$$

Intuitively, we may think of $\delta(x - x_i)$ as a very peak-shaped function $x = x_i$ with integral 1, roughly visualized in Figure 46. Because of (143), let $w_i = \delta(x - x_i)$ be weighting functions in the method of weighted reand (141) becomes equivalent to (142).

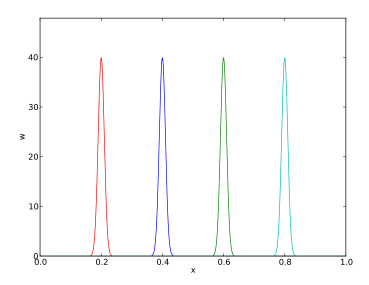


Figure 46: Approximation of delta functions by narrow Gaussian func

The subdomain collocation method. The idea of this approach is mand the integral of R to vanish over N+1 subdomains Ω_i of Ω :

$$\int_{\Omega_i} R \, \mathrm{d}x = 0, \quad i \in \mathcal{I}_s \, .$$

This statement can also be expressed as a weighted residual method

$$\int_{\Omega} Rw_i \, dx = 0, \quad i \in \mathcal{I}_s,$$

where $w_i = 1$ for $x \in \Omega_i$ and $w_i = 0$ otherwise.

1.9 Examples on using the principles

et us now apply global basis functions to illustrate the principles for minimizing $\dot{}$.

'he model problem. We consider the differential equation problem

$$-u''(x) = f(x), \quad x \in \Omega = [0, L], \quad u(0) = 0, \ u(L) = 0.$$
 (146)

lasis functions. Our choice of basis functions ψ_i for V is

$$\psi_i(x) = \sin\left((i+1)\pi\frac{x}{L}\right), \quad i \in \mathcal{I}_s.$$
 (147)

An important property of these functions is that $\psi_i(0) = \psi_i(L) = 0$, which leans that the boundary conditions on u are fulfilled:

$$u(0) = \sum_{j} c_j \psi_j(0) = 0, \quad u(L) = \sum_{j} c_j \psi_j(L) = 0.$$

nother nice property is that the chosen sine functions are orthogonal on Ω :

$$\int_{0}^{L} \sin\left((i+1)\pi\frac{x}{L}\right) \sin\left((j+1)\pi\frac{x}{L}\right) dx = \begin{cases} \frac{1}{2}L & i=j\\ 0, & i\neq j \end{cases}$$
 (148)

rovided i and j are integers.

'he residual. We can readily calculate the following explicit expression for ne residual:

$$R(x; c_0, \dots, c_N) = u''(x) + f(x),$$

$$= \frac{d^2}{dx^2} \left(\sum_{j \in \mathcal{I}_s} c_j \psi_j(x) \right) + f(x),$$

$$= \sum_{j \in \mathcal{I}_s} c_j \psi_j''(x) + f(x).$$
(149)

'he least squares method. The equations (137) in the least squares method equire an expression for $\partial R/\partial c_i$. We have

$$\frac{\partial R}{\partial c_i} = \frac{\partial}{\partial c_i} \left(\sum_{j \in \mathcal{I}_s} c_j \psi_j''(x) + f(x) \right) = \sum_{j \in \mathcal{I}_s} \frac{\partial c_j}{\partial c_i} \psi_j''(x) = \psi_i''(x) \,. \tag{150}$$

he governing equations for $\{c_i\}_{i\in\mathcal{I}_s}$ are then

$$\left(\sum_{j} c_{j} \psi_{j}^{"} + f, \psi_{i}^{"}\right) = 0, \quad i \in \mathcal{I}_{s},$$

which can be rearranged as

$$\sum_{j \in \mathcal{I}_s} (\psi_i'', \psi_j'') c_j = -(f, \psi_i''), \quad i \in \mathcal{I}_s.$$

This is nothing but a linear system

$$\sum_{j \in \mathcal{I}_s} A_{i,j} c_j = b_i, \quad i \in \mathcal{I}_s,$$

with

$$A_{i,j} = (\psi_i'', \psi_j'')$$

$$= \pi^4 (i+1)^2 (j+1)^2 L^{-4} \int_0^L \sin\left((i+1)\pi \frac{x}{L}\right) \sin\left((j+1)\pi \frac{x}{L}\right)$$

$$= \begin{cases} \frac{1}{2} L^{-3} \pi^4 (i+1)^4 & i=j\\ 0, & i\neq j \end{cases}$$

$$b_i = -(f, \psi_i'') = (i+1)^2 \pi^2 L^{-2} \int_0^L f(x) \sin\left((i+1)\pi \frac{x}{L}\right) dx$$

Since the coefficient matrix is diagonal we can easily solve for

$$c_i = \frac{2L}{\pi^2(i+1)^2} \int_0^L f(x) \sin((i+1)\pi \frac{x}{L}) dx.$$

With the special choice of f(x) = 2 can be calculated in sympy by

```
from sympy import *
import sys

i, j = symbols('i j', integer=True)
x, L = symbols('x L')
f = 2
a = 2*L/(pi**2*(i+1)**2)
c_i = a*integrate(f*sin((i+1)*pi*x/L), (x, 0, L))
c_i = simplify(c_i)
print c_i
```

The answer becomes

$$c_i = 4 \frac{L^2 \left(\left(-1 \right)^i + 1 \right)}{\pi^3 \left(i^3 + 3i^2 + 3i + 1 \right)}$$

Now, $1+(-1)^i=0$ for i odd, so only the coefficients with even index are Introducing i=2k for $k=0,\ldots,N/2$ to count the relevant indices (for k goes to (N-1)/2), we get the solution

$$u(x) = \sum_{k=0}^{N/2} \frac{8L^2}{\pi^3 (2k+1)^3} \sin\left((2k+1)\pi \frac{x}{L}\right). \tag{156}$$

he coefficients decay very fast: $c_2 = c_0/27$, $c_4 = c_0/125$. The solution will perefore be dominated by the first term.

$$u(x) \approx \frac{8L^2}{\pi^3} \sin\left(\pi \frac{x}{L}\right)$$
.

'he Galerkin method. The Galerkin principle (138) applied to (146) consists f inserting our special residual (149) in (138)

$$(u'' + f, v) = 0, \quad \forall v \in V,$$

r

$$(u'', v) = -(f, v), \quad \forall v \in V.$$

$$(157)$$

his is the variational formulation, based on the Galerkin principle, of our ifferential equation. The $\forall v \in V$ requirement is equivalent to demanding the quation (u'', v) = -(f, v) to be fulfilled for all basis functions $v = \psi_i$, $i \in \mathcal{I}_s$, $v \in (138)$ and $v \in$

$$\left(\sum_{j\in\mathcal{I}_s} c_j \psi_j'', \psi_i\right) = -(f, \psi_i), \quad i \in \mathcal{I}_s.$$
(158)

his equation can be rearranged to a form that explicitly shows that we get a near system for the unknowns $\{c_i\}_{i\in\mathcal{T}}$:

$$\sum_{j \in \mathcal{I}_s} (\psi_i, \psi_j'') c_j = (f, \psi_i), \quad i \in \mathcal{I}_s.$$
(159)

or the particular choice of the basis functions (147) we get in fact the same near system as in the least squares method because $\psi'' = -(i+1)^2 \pi^2 L^{-2} \psi$.

The collocation method. For the collocation method (142) we need to decide pon a set of N+1 collocation points in Ω . A simple choice is to use uniformly baced points: $x_i = i\Delta x$, where $\Delta x = L/N$ in our case $(N \ge 1)$. However, these oints lead to at least two rows in the matrix consisting of zeros (since $\psi_i(x_0) = 0$ and $\psi_i(x_N) = 0$), thereby making the matrix singular and non-invertible. This process us to choose some other collocation points, e.g., random points or points niformly distributed in the interior of Ω . Demanding the residual to vanish at ness points leads, in our model problem (146), to the equations

$$-\sum_{j\in\mathcal{I}_s} c_j \psi_j''(x_i) = f(x_i), \quad i\in\mathcal{I}_s,$$
(160)

hich is seen to be a linear system with entries

$$A_{i,j} = -\psi_j''(x_i) = (j+1)^2 \pi^2 L^{-2} \sin\left((j+1)\pi \frac{x_i}{L}\right),$$

in the coefficient matrix and entries $b_i = 2$ for the right-hand side f(x) = 2).

The special case of N=0 can sometimes be of interest. A natural of then the midpoint $x_0=L/2$ of the domain, resulting in $A_{0,0}=-\psi_0''(x_0)=f(x_0)=2$, and hence $c_0=2L^2/\pi^2$.

Comparison. In the present model problem, with f(x) = 2, the exact is u(x) = x(L-x), while for N=0 the Galerkin and least squares result in $u(x) = 8L^2\pi^{-3}\sin(\pi x/L)$ and the collocation method leads to $2L^2\pi^{-2}\sin(\pi x/L)$. We can quickly use sympy to verify that the maximu occurs at the midpoint x = L/2 and find what the errors are:

```
>>> import sympy as sp
>>> # Computing with Dirichlet conditions: -u''=2 and sines
>>> x, L = sp.symbols('x L')
>>> e_Galerkin = x*(L-x) - 8*L**2*sp.pi**(-3)*sp.sin(sp.pi*x/L)
>>> e_colloc = x*(L-x) - 2*L**2*sp.pi**(-2)*sp.sin(sp.pi*x/L)
>>> # Verify max error for x=L/2
>>> dedx_Galerkin = sp.diff(e_Galerkin, x)
>>> dedx_Galerkin.subs(x, L/2)
0
>>> dedx_colloc = sp.diff(e_colloc, x)
>>> dedx_colloc.subs(x, L/2)
0

# Compute max error: x=L/2, evaluate numerical, and simplify
>>> sp.simplify(e_Galerkin.subs(x, L/2).evalf(n=3))
-0.00812*L**2
>>> sp.simplify(e_colloc.subs(x, L/2).evalf(n=3))
0.0473*L**2
```

The error in the collocation method is about 6 times larger than the error Galerkin or least squares method.

11.10 Integration by parts

A problem arises if we want to apply popular finite element functions our model problem (146) by the standard least squares, Galerkin, or col methods: the piecewise polynomials $\psi_i(x)$ have discontinuous derivathe cell boundaries which makes it problematic to compute the second derivative. This fact actually makes the least squares and collocation method suitable for finite element approximation of the unknown function. (By rethe equation -u'' = f as a system of two first-order equations, u' = v and f, the least squares method can be applied. Also, differentiating disconfunctions can actually be handled by distribution theory in mathematic Galerkin method and the method of weighted residuals can, however, be

ogether with finite element basis functions if we use *integration by parts* as a leans for transforming a second-order derivative to a first-order one.

Consider the model problem (146) and its Galerkin formulation

$$-(u'', v) = (f, v) \quad \forall v \in V.$$

sing integration by parts in the Galerkin method, we can move a derivative of onto v:

$$\int_0^L u''(x)v(x) dx = -\int_0^L u'(x)v'(x) dx + [vu']_0^L$$
$$= -\int_0^L u'(x)v'(x) dx + u'(L)v(L) - u'(0)v(0). \tag{161}$$

sually, one integrates the problem at the stage where the u and v functions iter the formulation. Alternatively, but less common, we can integrate by parts i the expressions for the matrix entries:

$$\int_{0}^{L} \psi_{i}(x)\psi_{j}''(x) dx = -\int_{0}^{L} \psi_{i}'(x)\psi_{j}'(x)dx + [\psi_{i}\psi_{j}']_{0}^{L}$$

$$= -\int_{0}^{L} \psi_{i}'(x)\psi_{j}'(x) dx + \psi_{i}(L)\psi_{j}'(L) - \psi_{i}(0)\psi_{j}'(0). \quad (162)$$

itegration by parts serves to reduce the order of the derivatives and to make the befficient matrix symmetric since $(\psi_i', \psi_j') = (\psi_i', \psi_j')$. The symmetry property epends on the type of terms that enter the differential equation. As will be seen ter in Section 15, integration by parts also provides a method for implementing oundary conditions involving u'.

With the choice (147) of basis functions we see that the "boundary terms" $_i(L)\psi_j'(L)$ and $\psi_i(0)\psi_j'(0)$ vanish since $\psi_i(0)=\psi_i(L)=0$.

Veak form. Since the variational formulation after integration by parts make eaker demands on the differentiability of u and the basis functions ψ_i , the resulting integral formulation is referred to as a weak form of the differential equation roblem. The original variational formulation with second-order derivatives, or ne differential equation problem with second-order derivative, is then the strong prim, with stronger requirements on the differentiability of the functions.

For differential equations with second-order derivatives, expressed as variaonal formulations and solved by finite element methods, we will always perform itegration by parts to arrive at expressions involving only first-order derivatives.

1.11 Boundary function

o far we have assumed zero Dirichlet boundary conditions, typically u(0) = (L) = 0, and we have demanded that $\psi_i(0) = \psi_i(L) = 0$ for $i \in \mathcal{I}_s$. What

about a boundary condition like $u(L) = D \neq 0$? This condition immfaces a problem: $u = \sum_{i} c_{i} \varphi_{i}(L) = 0$ since all $\varphi_{i}(L) = 0$.

A boundary condition of the form u(L) = D can be implemented by a ing that all $\psi_i(L) = 0$, but adding a boundary function B(x) with the boundary value, B(L) = D, to the expansion for u:

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x).$$

This u gets the right value at x = L:

$$u(L) = B(L) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(L) = B(L) = D.$$

The idea is that for any boundary where u is known we demand ψ_i to and construct a function B(x) to attain the boundary value of u. The no restrictions how B(x) varies with x in the interior of the domain, variation needs to be constructed in some way.

For example, with u(0) = 0 and u(L) = D, we can choose B(x) =since this form ensures that B(x) fulfills the boundary conditions: B(0) B(L) = D. The unknown function is then sought on the form

$$u(x) = \frac{x}{L}D + \sum_{j \in \mathcal{I}_c} c_j \psi_j(x),$$

with $\psi_i(0) = \psi_i(L) = 0$.

The B(x) function can be chosen in many ways as long as its boundar are correct. For example, $B(x) = D(x/L)^p$ for any power p will work fin above example.

As another example, consider a domain $\Omega = [a, b]$ where the beconditions are $u(a) = U_a$ and $u(b) = U_b$. A class of possible B(x) funct

$$B(x) = U_a + \frac{U_b - U_a}{(b-a)^p} (x-a)^p, \quad p > 0.$$

Real applications will most likely use the simplest version, p = 1, but he a p parameter was included to demonstrate the ambiguity in the construB(x).

Summary.

The general procedure of incorporating Dirichlet boundary conditions as follows. Let $\partial \Omega_E$ be the part(s) of the boundary $\partial \Omega$ of the doma where u is specified. Set $\psi_i = 0$ at the points in $\partial \Omega_E$ and seek u as

$$u(x) = B(x) + \sum_{j \in \mathcal{T}_0} c_j \psi_j(x),$$

temark. With the B(x) term, u does not in general lie in $V = \operatorname{span} \{\psi_0, \dots, \psi_N\}$ nymore. Moreover, when a prescribed value of u at the boundary, say $u(a) = U_a$ different from zero, it does not make sense to say that u lies in a vector space, ecause this space does not obey the requirements of addition and scalar mulplication. For example, 2u does not lie in the space since its boundary value $2U_a$, which is incorrect. It only makes sense to split u in two parts, as done bove, and have the unknown part $\sum_i c_j \psi_j$ in a proper function space.

1.12 Abstract notation for variational formulations

We have seen that variational formulations end up with a formula involving u and such as (u',v') and a formula involving v and known functions, such as (f,v). Widely used notation is to introduce an abstract variational statement written s a(u,v) = L(v), where a(u,v) is a so-called bilinear form involving all the erms that contain both the test and trial function, while L(v) is a linear form ontaining all the terms without the trial function. For example, the statement

$$\int_{\Omega} u'v' \, dx = \int_{\Omega} fv \, dx \quad \text{or} \quad (u', v') = (f, v) \quad \forall v \in V$$

an be written in abstract form: find u such that

$$a(u, v) = L(v) \quad \forall v \in V,$$

here we have the definitions

$$a(u, v) = (u', v'), L(v) = (f, v).$$

The term linear means that $L(\alpha_1v_1 + \alpha_2v_2) = \alpha_1L(v_1) + \alpha_2L(v_2)$ for two est functions v_1 and v_2 , and scalar parameters α_1 and α_2 . Similarly, the term linear means that a(u,v) is linear in both its arguments:

$$a(\alpha_1 u_1 + \alpha_2 u_2, v) = \alpha_1 a(u_1, v) + \alpha_2 a(u_2, v),$$

$$a(u, \alpha_1 v_1 + \alpha_2 v_2) = \alpha_1 a(u, v_1) + \alpha_2 a(u, v_2).$$

1 nonlinear problems these linearity properties do not hold in general and the betract notation is then F(u; v) = 0.

The matrix system associated with a(u,v) = L(v) can also be written in an betract form by inserting $v = \psi_i$ and $u = \sum_j c_j \psi_j$ in a(u,v) = L(v). Using the near properties, we get

$$\sum_{j \in \mathcal{I}_s} a(\psi_j, \psi_i) c_j = L(\psi_i), \quad i \in \mathcal{I}_s,$$

hich is a linear system

$$\sum_{j \in \mathcal{I}_s} A_{i,j} c_j = b_i, \quad i \in \mathcal{I}_s,$$

where

$$A_{i,j} = a(\psi_i, \psi_i), \quad b_i = L(\psi_i).$$

In many problems, a(u, v) is symmetric such that $a(\psi_j, \psi_i) = a(\psi_i, \psi_j)$. cases the coefficient matrix becomes symmetric, $A_{i,j} = A_{j,i}$, a property t simplify solution algorithms for linear systems and make them more s addition to saving memory and computations.

The abstract notation a(u,v) = L(v) for linear differential equation p is much used in the literature and in description of finite element s (in particular the FEniCS²¹ documentation). We shall frequently sun variational forms using this notation.

11.13 Variational problems and optimization of funals

If a(u,v) = a(v,u), it can be shown that the variational statement

$$a(u, v) = L(v) \quad \forall v \in V,$$

is equivalent to minimizing the functional

$$F(v) = \frac{1}{2}a(v,v) - L(v)$$

over all functions $v \in V$. That is,

$$F(u) \le F(v) \quad \forall v \in V.$$

Inserting a $v=\sum_j c_j \psi_j$ turns minimization of F(v) into minimization quadratic function

$$\bar{F}(c_0, \dots, c_N) = \sum_{j \in \mathcal{I}_s} \sum_{i \in \mathcal{I}_s} a(\psi_i, \psi_j) c_i c_j - \sum_{j \in \mathcal{I}_s} L(\psi_j) c_j$$

of N+1 parameters.

Minimization of F implies

$$\frac{\partial \bar{F}}{\partial c_i} = 0, \quad i \in \mathcal{I}_s.$$

After some algebra one finds

$$\sum_{j} j \in \mathcal{I}_s a(\psi_i, \psi_j) c_j = L(\psi_i), \quad i \in \mathcal{I}_s,$$

 $^{^{21} {}m http://fenicsproject.org}$

hich is the same system as that arising from a(u, v) = L(v).

Many traditional applications of the finite element method, especially in solid nechanics and structural analysis, start with formulating F(v) from physical rinciples, such as minimization of energy, and then proceeds with deriving (u, v) = L(v), which is the equation usually desired in implementations.

2 Examples on variational formulations

he following sections derive variational formulations for some prototype differntial equations in 1D, and demonstrate how we with ease can handle variable pefficients, mixed Dirichlet and Neumann boundary conditions, first-order erivatives, and nonlinearities.

2.1 Variable coefficient

onsider the problem

$$-\frac{d}{dx}\left(\alpha(x)\frac{du}{dx}\right) = f(x), \quad x \in \Omega = [0, L], \ u(0) = C, \ u(L) = D. \tag{166}$$

here are two new features of this problem compared with previous examples: variable coefficient a(x) and nonzero Dirichlet conditions at both boundary oints.

Let us first deal with the boundary conditions. We seek

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_i(x),$$

ith $\psi_i(0) = \psi_i(L) = 0$ for $i \in \mathcal{I}_s$. The function B(x) must then fulfill B(0) = C and B(L) = D. How B varies in between x = 0 and x = L is not of importance. The possible choice is

$$B(x) = C + \frac{1}{L}(D - C)x,$$

hich follows from (164) with p = 1.

We seek $(u - B) \in V$. As usual,

$$V = \operatorname{span}\{\psi_0, \dots, \psi_N\},\$$

ut the two Dirichlet boundary conditions demand that

$$\psi_i(0) = \psi_i(L) = 0, \quad i \in \mathcal{I}_s.$$

ote that any $v \in V$ has the property v(0) = v(L) = 0.

The residual arises by inserting our u in the differential equation:

$$R = -\frac{d}{dx} \left(\alpha \frac{du}{dx} \right) - f.$$

Galerkin's method is

$$(R, v) = 0, \quad \forall v \in V,$$

or written with explicit integrals,

$$\int_{\Omega} \left(\frac{d}{dx} \left(\alpha \frac{du}{dx} \right) - f \right) v \, dx = 0, \quad \forall v \in V.$$

We proceed with integration by parts to lower the derivative from second order:

$$-\int_{\Omega} \frac{d}{dx} \left(\alpha(x) \frac{du}{dx} \right) v \, dx = \int_{\Omega} \alpha(x) \frac{du}{dx} \frac{dv}{dx} \, dx - \left[\alpha \frac{du}{dx} v \right]_{0}^{L}.$$

The boundary term vanishes since v(0) = v(L) = 0. The variational tion is then

$$\int_{\Omega} \alpha(x) \frac{du}{dx} \frac{dv}{dx} dx = \int_{\Omega} f(x)v dx, \quad \forall v \in V.$$

The variational formulation can alternatively be written in a more compa

$$(\alpha u', v') = (f, v), \quad \forall v \in V.$$

The corresponding abstract notation reads

$$a(u, v) = L(v) \quad \forall v \in V,$$

with

$$a(u, v) = (\alpha u', v'), \quad L(v) = (f, v).$$

Note that the a in the notation $a(\cdot, \cdot)$ is not to be mixed with the coefficient a(x) in the differential equation.

We may insert $u = B + \sum_{j} c_{j} \psi_{j}$ and $v = \psi_{i}$ to derive the linear sys

$$(\alpha B' + \alpha \sum_{j \in \mathcal{I}_s} c_j \psi'_j, \psi'_i) = (f, \psi_i), \quad i \in \mathcal{I}_s.$$

Isolating everything with the c_j coefficients on the left-hand side and al terms on the right-hand side gives

$$\sum_{j \in \mathcal{I}_s} (\alpha \psi_j', \psi_i') c_j = (f, \psi_i) + (a(D - C)L^{-1}, \psi_i'), \quad i \in \mathcal{I}_s.$$

This is nothing but a linear system $\sum_{i} A_{i,j} c_j = b_i$ with

$$\mathbf{1}_{i,j} = (a\psi'_j, \psi'_i) = \int_{\Omega} \alpha(x)\psi'_j(x), \psi'_i(x) \, dx,
b_i = (f, \psi_i) + (a(D - C)L^{-1}, \psi'_i) = \int_{\Omega} \left(f(x)\psi_i(x) + \alpha(x) \frac{D - C}{L} \psi'_i(x) \right) \, dx.$$

2.2 First-order derivative in the equation and boundary condition

he next problem to formulate in variational form reads

$$-u''(x) + bu'(x) = f(x), \quad x \in \Omega = [0, L], \ u(0) = C, \ u'(L) = E. \tag{167}$$

he new features are a first-order derivative u' in the equation and the boundary ondition involving the derivative: u'(L) = E. Since we have a Dirichlet condition t x = 0, we must force $\psi_i(0) = 0$ and use a boundary function to take care of ne condition u(0) = C. Because there is no Dirichlet condition on x = L we do ot make any requirements to $\psi_i(L)$. The simplest possible choice of B(x) is f(x) = C.

The expansion for u becomes

$$u = C + \sum_{j \in \mathcal{I}_s} c_j \psi_i(x) .$$

The variational formulation arises from multiplying the equation by a test unction $v \in V$ and integrating over Ω :

$$(-u'' + bu' - f, v) = 0, \quad \forall v \in V$$

We apply integration by parts to the u''v term only. Although we could also itegrate u'v by parts, this is not common. The result becomes

$$(u' + bu', v') = (f, v) + [u'v]_0^L, \quad \forall v \in V.$$

ow, v(0) = 0 so

$$[u'v]_0^L = u'(L)v(L) = Ev(L),$$

ecause u'(L) = E. Integration by parts allows us to take care of the Neumann ondition in the boundary term.

Natural and essential boundary conditions.

Omitting a boundary term like $[u'v]_0^L$ implies that we actually impose the condition u'=0 unless there is a Dirichlet condition (i.e., v=0) at that point! This result has great practical consequences, because it is easy to forget the boundary term, and this mistake may implicitly set a boundary

condition! Since homogeneous Neumann conditions can be incorporated without doing anything, and non-homogeneous Neumann conditions just be inserted in the boundary term, such conditions are known as no boundary conditions. Dirichlet conditions requires more essential stee the mathematical formulation, such as forcing all $\varphi_i = 0$ on the bour and constructing a B(x), and are therefore known as essential bour conditions.

The final variational form reads

$$(u', v') + (bu', v) = (f, v) + Ev(L), \quad \forall v \in V.$$

In the abstract notation we have

$$a(u, v) = L(v) \quad \forall v \in V,$$

with the particular formulas

$$a(u, v) = (u', v') + (bu', v), \quad L(v) = (f, v) + Ev(L).$$

The associated linear system is derived by inserting $u = B + \sum_j c_j$ replacing v by ψ_i for $i \in \mathcal{I}_s$. Some algebra results in

$$\sum_{j \in \mathcal{I}_s} \underbrace{((\psi'_j, \psi'_i) + (b\psi'_j, \psi_i))}_{A_{i,i}} c_j = \underbrace{(f, \psi_i) + E\psi_i(L)}_{b_i}.$$

Observe that in this problem, the coefficient matrix is not symmetric, of the term

$$(b\psi'_j, \psi_i) = \int_{\Omega} b\psi'_j \psi_i \, \mathrm{d}x \neq \int_{\Omega} b\psi'_i \psi_j \, \mathrm{d}x = (\psi'_i, b\psi_j) \,.$$

12.3 Nonlinear coefficient

Finally, we show that the techniques used above to derive variations also apply to nonlinear differential equation problems as well. Here is a problem with a nonlinear coefficient and right-hand side:

$$-(\alpha(u)u')' = f(u), \quad x \in [0, L], \ u(0) = 0, \ u'(L) = E.$$

Our space V has basis $\{\psi_i\}_{i\in\mathcal{I}_s}$, and because of the condition u(0)=0, require $\psi_i(0)=0$, $i\in\mathcal{I}_s$.

Galerkin's method is about inserting the approximate u, multiply differential equation by $v \in V$, and integrate,

$$-\int_0^L \frac{d}{dx} \left(\alpha(u) \frac{du}{dx} \right) v \, dx = \int_0^L f(u) v \, dx \quad \forall v \in V.$$

he integration by parts does not differ from the case where we have $\alpha(x)$ instead $\alpha(u)$:

$$\int_0^L \alpha(u) \frac{du}{dx} \frac{dv}{dx} dx = \int_0^L f(u)v dx + [\alpha(u)vu']_0^L \quad \forall v \in V.$$

he term $\alpha(u(0))v(0)u'(0) = 0$ since v(0). The other term, $\alpha(u(L))v(L)u'(L)$, is sed to impose the other boundary condition u'(L) = E, resulting in

$$\int_0^L \alpha(u) \frac{du}{dx} \frac{dv}{dx} v \, dx = \int_0^L f(u)v \, dx + \alpha(u(L))v(L)E \quad \forall v \in V,$$

r alternatively written more compactly as

$$(\alpha(u)u', v') = (f(u), v) + \alpha(L)v(L)E \quad \forall v \in V.$$

ince the problem is nonlinear, we cannot identify a bilinear form a(u, v) and a near form L(v). An abstract notation is typically find u such that

$$F(u; v) = 0 \quad \forall v \in V,$$

ith

$$F(u; v) = (a(u)u', v') - (f(u), v) - a(L)v(L)E.$$

By inserting $u = \sum_j c_j \psi_j$ we get a nonlinear system of algebraic equations or the unknowns c_i , $i \in \mathcal{I}_s$. Such systems must be solved by constructing a equence of linear systems whose solutions hopefully converge to the solution f the nonlinear system. Frequently applied methods are Picard iteration and ewton's method.

2.4 Computing with Dirichlet and Neumann conditions

et us perform the necessary calculations to solve

$$-u''(x) = 2$$
, $x \in \Omega = [0, 1]$, $u'(0) = C$, $u(1) = D$,

sing a global polynomial basis $\psi_i \sim x^i$. The requirements on ψ_i is that $\psi_i(1) = 0$, ecause u is specified at x = 1, so a proper set of polynomial basis functions can e

$$\psi_i(x) = (1-x)^{i+1}, \quad i \in \mathcal{I}_s.$$

suitable B(x) function to handle the boundary condition u(1) = D is B(x) = Dx. The variational formulation becomes

$$(u', v') = (2, v) - Cv(0) \quad \forall v \in V.$$

he entries in the linear system are then

$$A_{i,j} = (\psi_j, \psi_i) = \int_0^1 \psi_i'(x)\psi_j'(x) \, \mathrm{d}x = \int_0^1 (i+1)(j+1)(1-x)^{i+j} \, \mathrm{d}x = \int_0^1 (2\psi_i(x) - (D, \psi_i') - C\psi_i(0))$$

$$= \int_0^1 (2\psi_i(x) - D\psi_i'(x)) \, \mathrm{d}x - C\psi_i(0)$$

$$= \int_0^1 (2(1-x)^{i+1} - D(i+1)(1-x)^i) \, \mathrm{d}x - C\psi_i(0)$$

$$= \frac{2 - (2+i)(D+C)}{i+2}.$$

With N=1 the global matrix system is

$$\begin{pmatrix} 1 & 1 \\ 1 & 4/3 \end{pmatrix} \begin{pmatrix} c_0 \\ c_1 \end{pmatrix} = \begin{pmatrix} -C+D+1 \\ 2/3-C+D \end{pmatrix}$$

The solution becomes $c_0 = -C + D + 2$ and $c_1 = -1$, resulting in

$$u(x) = 1 - x^2 + D + C(x - 1),$$

The exact solution is found by. integrating twice and applying the beconditions, either by hand or using sympy as shown in Section 11.2. It that the numerical solution coincides with the exact one. This result expected because if $(u_e - B) \in V$, $u = u_e$, as proved next.

12.5 When the numerical method is exact

We have some variational formulation: find $(u-B) \in V$ such that a $L(u) \, \forall V$. The exact solution also fulfills $a(u_e,v) = L(v)$, but normally (lies in a much larger (infinite-dimensional) space. Suppose, neverthele $u_e = B + E$, where $E \in V$. That is, apart from Dirichlet conditions, u_e our finite-dimensional space V we use to compute u. Writing also u on t form u = B + F, we have

$$a(B+E,v) = L(v) \quad \forall v \in V,$$

 $a(B+F,v) = L(v) \quad \forall v \in V.$

Subtracting the equations show that a(E-F,v)=0 for all $v\in V$, and t E-F=0 and $u=u_{\rm e}$.

The case treated in Section 12.4 is of the type where $u_e - B$ is a quantum function that is 0 at x = 1, and therefore $(u_e - B) \in V$, and the method the exact solution.

3 Computing with finite elements

he purpose of this section is to demonstrate in detail how the finite element lethod can the be applied to the model problem

$$-u''(x) = 2$$
, $x \in (0, L)$, $u(0) = u(L) = 0$,

ith variational formulation

$$(u', v') = (2, v) \quad \forall v \in V.$$

he variational formulation is derived in Section 11.10.

3.1 Finite element mesh and basis functions

/e introduce a finite element mesh with N_e cells, all with length h, and number ne cells from left to right. global nodes. Choosing P1 elements, there are two odes per cell, and the coordinates of the nodes become

$$x_i = ih, \quad h = L/N_e, \quad i = 0, \dots, N_n = N_e + 1,$$

rovided we number the nodes from left to right.

Each of the nodes, i, is associated a finite element basis function $\varphi_i(x)$. When pproximating a given function f by a finite element function u, we expand using finite element basis functions associated with all nodes in the mesh, e., $N=N_n$. However, when solving differential equations we will often have $\ell < N_n$ because of Dirichlet boundary conditions. Why this is the case will ow be explained in detail.

In our case with homogeneous Dirichlet boundary conditions we do not need ny boundary function B(x) and can work with the expansion

$$u(x) = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x). \tag{170}$$

ecause of the boundary conditions, we must demand $\psi_i(0) = \psi_i(L) = 0$, $i \in \mathcal{I}_s$. Then ψ_i , $i = 0, \ldots, N$, is to be selected among the finite element basis functions j, $i = 0, \ldots, N_n$, we have to avoid using φ_j functions that do not vanish at 0 = 0 and $x_{N_n} = L$. However, all φ_j vanish at these two nodes for $j = 1, \ldots, N_n$. Inly basis functions associated with the end nodes, φ_0 and φ_{N_n} , violate the oundary conditions of our differential equation. Therefore, we select the basis motions φ_i to be the set of finite element basis functions associated with all ne interior nodes in the mesh:

$$\psi_i = \varphi_{i+1}, \quad i = 0, \dots, N.$$

fere, $N = N_n - 2$.

In the general case, the nodes are not necessarily numbered from left to right, by we introduce a mapping from the node numbering, or more precisely the egree of freedom numbering, to the numbering of the unknowns in the final

equation system. These unknowns take on the numbers $0, \ldots, N$. Us number j in the linear system corresponds to degree of freedom numb $j \in \mathcal{I}_s$. We can then write

$$\psi_i = \varphi_{\nu(i)}, \quad i = 0, \dots, N.$$

With a regular numbering as in the present example, $\nu(j)=j+1,\,j=1,.$ $N_n-2.$

13.2 Computation in the global physical domain

We shall first perform a computation in the x coordinate system becautegrals can be easily computed here by simple, visual, geometric considerable that the coordinate system is called a global approach since we work in the x coordinate system compute integrals on the global domain [0, L].

The entries in the coefficient matrix and right-hand side are

$$A_{i,j} = \int_0^L \psi_i'(x)\psi_j'(x) dx, \quad b_i = \int_0^L 2\psi_i(x) dx, \quad i, j \in \mathcal{I}_s.$$

Expressed in terms of finite element basis functions φ_i we get the alterespressions

$$A_{i,j} = \int_0^L \varphi'_{i+1}(x)\varphi'_{j+1}(x) dx, \quad b_i = \int_0^L 2\varphi_{i+1}(x) dx, \quad i, j \in \mathcal{I}_s$$

For the following calculations the subscripts on the finite element basis for are more conveniently written as i and j instead of i+1 and j+1 notation becomes

$$A_{i-1,j-1} = \int_0^L \varphi_i'(x)\varphi_j'(x) dx, \quad b_{i-1} = \int_0^L 2\varphi_i(x) dx,$$

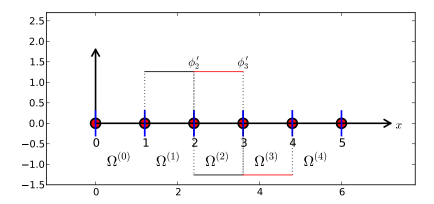
where the i and j indices run as $i, j = 1, ..., N_n - 1 = N + 1$.

The $\varphi_i(x)$ function is a hat function with peak at $x = x_i$ and a linear v in $[x_{i-1}, x_i]$ and $[x_i, x_{i+1}]$. The derivative is 1/h to the left of x_i and the right, or more formally,

$$\varphi_i'(x) = \begin{cases} 0, & x < x_{i-1}, \\ h^{-1}, & x_{i-1} \le x < x_i, \\ -h^{-1}, & x_i \le x < x_{i+1}, \\ 0, & x \ge x_{i+1} \end{cases}$$

Figure 47 shows $\varphi'_1(x)$ and $\varphi'_2(x)$.

We realize that φ'_i and $\bar{\varphi}'_j$ has no overlap, and hence their productions, unless i and j are nodes belonging to the same cell. The only contributions to the coefficient matrix are therefore



igure 47: Illustration of the derivative of piecewise linear basis functions ssociated with nodes in cell 2.

$$A_{i-1,i-2} = \int_0^L \varphi_i'(x)\varphi_{i-1}'(x) \, dx,$$

$$A_{i-1,i-1} = \int_0^L \varphi_i'(x)^2 \, dx,$$

$$A_{i-1,i} = \int_0^L \varphi_i'(x)\varphi_{i+1}'(x) \, dx,$$

or $i = 1, ..., N_n - 1$, but for $i = 1, A_{i-1,i-2}$ is not defined, and for $i = N_n - 1$, i-1,i is not defined.

We see that $\varphi'_{i-1}(x)$ and $\varphi'_i(x)$ have overlap of one cell $\Omega^{(i-1)} = [x_{i-1}, x_i]$ nd that their product then is $-1/h^2$. The integrand is constant and therefore $i-1,i-2=-h^{-2}h=-h^{-1}$. A similar reasoning can be applied to $A_{i-1,i}$, which lso becomes $-h^{-1}$. The integral of $\varphi'_i(x)^2$ gets contributions from two cells, $f^{(i-1)} = [x_{i-1}, x_i]$ and $f^{(i)} = [x_i, x_{i+1}]$, but $f^{(i)} = [x_i, x_{i+1}]$ but $f^{(i)} = [x_i, x_{i+1}]$ but $f^{(i)} = [x_i, x_{i+1}]$.

The right-hand side involves an integral of $2\varphi_i(x)$, $i = 1, ..., N_n - 1$, which just the area under a hat function of height 1 and width 2h, i.e., equal to h. hence, $b_{i-1} = 2h$.

To summarize the linear system, we switch from i to i+1 such that we can rite

$$A_{i,i-1} = A_{i,i-1} = -h^{-1}, \quad A_{i,i} = 2h^{-1}, \quad b_i = 2h.$$

The equation system to be solved only involves the unknowns c_i for $i \in \mathcal{I}_s$. /ith our numbering of unknowns and nodes, we have that c_i equals $u(x_{i+1})$. he complete matrix system that takes the following form:

$$\frac{1}{h} \begin{pmatrix} 2 & -1 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\ -1 & 2 & -1 & \ddots & & & & \vdots \\ 0 & -1 & 2 & -1 & \ddots & & & \vdots \\ \vdots & \ddots & & \ddots & \ddots & \ddots & \ddots & \vdots \\ \vdots & & & 0 & -1 & 2 & -1 & \ddots & \vdots \\ \vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & & \ddots & \ddots & \ddots & \ddots & -1 \\ 0 & \cdots & \cdots & \cdots & \cdots & \cdots & 0 & -1 & 2 \end{pmatrix} \begin{pmatrix} c_0 \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N \end{pmatrix}$$

13.3 Comparison with a finite difference discretizat

A typical row in the matrix system can be written as

$$-\frac{1}{h}c_{i-1} + \frac{2}{h}c_i - \frac{1}{h}c_{i+1} = 2h.$$

Let us introduce the notation u_j for the value of u at node j: $u_j = u(x)$ we have the interpretation $u(x_j) = \sum_j c_j \varphi(x_j) = \sum_j c_j \delta_{ij} = c_j$. The us c_0, \ldots, c_N are u_1, \ldots, u_{N_n} . Shifting i with i+1 in (173) and inserting u_i we get

$$-\frac{1}{h}u_{i-1} + \frac{2}{h}u_i - \frac{1}{h}u_{i+1} = 2h,$$

A finite difference discretization of -u''(x) = 2 by a centered, secon finite difference approximation $u''(x_i) \approx [D_x D_x u]_i$ with $\Delta x = h$ yields

$$-\frac{u_{i-1}-2u_i+u_{i+1}}{h^2}=2,$$

which is, in fact, equivalent to (174) if (174) is divided by h. Theref finite difference and the finite element method are equivalent in this simproblem.

Sometimes a finite element method generates the finite difference ec on a uniform mesh, and sometimes the finite element method generates ec that are different. The differences are modest, but may influence the nu quality of the solution significantly, especially in time-dependent proble

13.4 Cellwise computations

We now employ the cell by cell computational procedure where an element and vector are calculated for each cell and assembled in the global linear Il integrals are mapped to the local reference coordinate system $X \in [-1,1]$.

1 the present case, the matrix entries contain derivatives with respect to x,

$$A_{i-1,j-1}^{(e)} = \int_{\Omega^{(e)}} \varphi_i'(x)\varphi_j'(x) \,\mathrm{d}x = \int_{-1}^1 \frac{d}{dx} \tilde{\varphi}_r(X) \frac{d}{dx} \tilde{\varphi}_s(X) \frac{h}{2} \,\mathrm{d}X,$$

here the global degree of freedom i is related to the local degree of freedom r irough i = q(e, r). Similarly, j = q(e, s). The local degrees of freedom run as s = 0, 1 for a P1 element.

'he integral for the element matrix. There are simple formulas for the asis functions $\tilde{\varphi}_r(X)$ as functions of X. However, we now need to find the erivative of $\tilde{\varphi}_r(X)$ with respect to x. Given

$$\tilde{\varphi}_0(X) = \frac{1}{2}(1 - X), \quad \tilde{\varphi}_1(X) = \frac{1}{2}(1 + X),$$

e can easily compute $d\tilde{\varphi}_r/dX$:

$$\frac{d\tilde{\varphi}_0}{dX} = -\frac{1}{2}, \quad \frac{d\tilde{\varphi}_1}{dX} = \frac{1}{2}.$$

rom the chain rule,

$$\frac{d\tilde{\varphi}_r}{dx} = \frac{d\tilde{\varphi}_r}{dX}\frac{dX}{dx} = \frac{2}{h}\frac{d\tilde{\varphi}_r}{dX}.$$
 (176)

he transformed integral is then

$$A_{i-1,j-1}^{(e)} = \int_{\Omega^{(e)}} \varphi_i'(x) \varphi_j'(x) \, \mathrm{d}x = \int_{-1}^1 \frac{2}{h} \frac{d\tilde{\varphi}_r}{dX} \frac{2}{h} \frac{d\tilde{\varphi}_s}{dX} \frac{h}{2} \, \mathrm{d}X.$$

'he integral for the element vector. The right-hand side is transformed coording to

$$b_{i-1}^{(e)} = \int_{\Omega^{(e)}} 2\varphi_i(x) \, \mathrm{d}x = \int_{-1}^1 2\tilde{\varphi}_r(X) \frac{h}{2} \, \mathrm{d}X, \quad i = q(e, r), \ r = 0, 1.$$

Petailed calculations of the element matrix and vector. Specifically or P1 elements we arrive at the following calculations for the element matrix atries:

$$\begin{split} \tilde{A}_{0,0}^{(e)} &= \int_{-1}^{1} \frac{2}{h} \left(-\frac{1}{2} \right) \frac{2}{h} \left(-\frac{1}{2} \right) \frac{2}{h} \, \mathrm{d}X = \frac{1}{h} \\ \tilde{A}_{0,1}^{(e)} &= \int_{-1}^{1} \frac{2}{h} \left(-\frac{1}{2} \right) \frac{2}{h} \left(\frac{1}{2} \right) \frac{2}{h} \, \mathrm{d}X = -\frac{1}{h} \\ \tilde{A}_{1,0}^{(e)} &= \int_{-1}^{1} \frac{2}{h} \left(\frac{1}{2} \right) \frac{2}{h} \left(-\frac{1}{2} \right) \frac{2}{h} \, \mathrm{d}X = -\frac{1}{h} \\ \tilde{A}_{1,1}^{(e)} &= \int_{-1}^{1} \frac{2}{h} \left(\frac{1}{2} \right) \frac{2}{h} \left(\frac{1}{2} \right) \frac{2}{h} \, \mathrm{d}X = \frac{1}{h} \end{split}$$

The element vector entries become

$$\tilde{b}_0^{(e)} = \int_{-1}^1 2\frac{1}{2}(1-X)\frac{h}{2} \, dX = h$$

$$\tilde{b}_1^{(e)} = \int_{-1}^1 2\frac{1}{2}(1+X)\frac{h}{2} \, dX = h.$$

Expressing these entries in matrix and vector notation, we have

$$\tilde{A}^{(e)} = \frac{1}{h} \left(\begin{array}{cc} 1 & -1 \\ -1 & 1 \end{array} \right), \quad \tilde{b}^{(e)} = h \left(\begin{array}{c} 1 \\ 1 \end{array} \right).$$

Contributions from the first and last cell. The first and last cell only one unknown and one basis function because of the Dirichlet be conditions at the first and last node. The element matrix therefore been 1×1 matrix and there is only one entry in the element vector. On cell $\psi_0 = \varphi_1$ is involved, corresponding to integration with $\tilde{\varphi}_1$. On cell $\psi_N = \varphi_{N_n-1}$ is involved, corresponding to integration with $\tilde{\varphi}_0$. We then special end-cell contributions

$$\tilde{A}^{(e)} = \frac{1}{h} \begin{pmatrix} 1 \end{pmatrix}, \quad \tilde{b}^{(e)} = h \begin{pmatrix} 1 \end{pmatrix},$$

for e = 0 and $e = N_e$. In these cells, we have only one degree of freed two as in the interior cells.

Assembly. The next step is to assemble the contributions from the cells. The assembly of an element matrix and vector into the global matright-hand side can be expressed as

$$A_{q(e,r),q(e,s)} = A_{q(e,r),q(e,s)} + \tilde{A}_{r,s}^{(e)}, \quad b_{q(e,r)} = b_{q(e,r)} + \tilde{b}_{r}^{(e)},$$

for r and s running over all local degrees of freedom in cell e.

To make the assembly algorithm more precise, it is convenient to Python data structures and a code snippet for carrying out all detail algorithm. For a mesh of four equal-sized P1 elements and L=2 we have

```
rertices = [0, 0.5, 1, 1.5, 2]
sells = [[0, 1], [1, 2], [2, 3], [3, 4]]
lof_map = [[0], [0, 1], [1, 2], [2]]
```

he total number of degrees of freedom is 3, being the function values at the iternal 3 nodes where u is unknown. In cell 0 we have global degree of freedom , the next cell has u unknown at its two nodes, which become global degrees of eedom 0 and 1, and so forth according to the dof_{map} list. The mathematical (e,r) quantity is nothing but the dof_{map} list.

Assume all element matrices are stored in a list Ae such that Ae[e][i,j] is $\tilde{b}_r^{(e)}$. A corresponding list for the element vectors is named be, where be[e][r] $\tilde{b}_r^{(e)}$. A Python code snippet illustrates all details of the assembly algorithm:

```
# A[i,j]: coefficient matrix, b[i]: right-hand side
for e in range(len(Ae)):
    for r in range(Ae[e].shape[0]):
        for s in range(Ae[e].shape[1]):
        A[dof_map[e,r],dof_map[e,s]] += Ae[e][i,j]
        b[dof_map[e,r]] += be[e][i,j]
```

The general case with N_e P1 elements of length h has

```
\[ n = N_e + 1 \]
rertices = [i*h for i in range(N_n)]
:ells = [[e, e+1] for e in range(N_e)]
lof_map = [[0]] + [[e-1, e] for i in range(1, N_e)] + [[N_n-2]]
```

Carrying out the assembly results in a linear system that is identical to (172), hich is not surprising since the procedures is mathematically equivalent to the alculations in the physical domain.

A fundamental problem with the matrix system we have assembled is that ne boundary conditions are not incorporated if u(0) or u(L) are different from ero. The next sections deals with this issue.

4 Boundary conditions: specified nonzero value

We have to take special actions to incorporate Dirichlet conditions, such as (L) = D, into the computational procedures. The present section outlines lternative, yet mathematically equivalent, methods.

4.1 General construction of a boundary function

1 Section 11.11 we introduce a boundary function B(x) to deal with nonzero irichlet boundary conditions for u. The construction of such a function is not lways trivial, especially not in multiple dimensions. However, a simple and eneral construction idea exists when the basis functions have the property

$$\varphi_i(x_j) = \delta_{ij}, \quad \delta_{ij} = \begin{cases} 1, & i = j, \\ 0, & i \neq j, \end{cases}$$

where x_j is a boundary point. Examples on such functions are the Lainterpolating polynomials and finite element functions.

Suppose now that u has Dirichlet boundary conditions at nodes with $i \in I_b$. For example, $I_b = \{0, N_n\}$ in a 1D mesh with node numbering f to right. Let U_i be the corresponding prescribed values of $u(x_i)$. We can general, use

$$B(x) = \sum_{j \in I_b} U_j \varphi_j(x) .$$

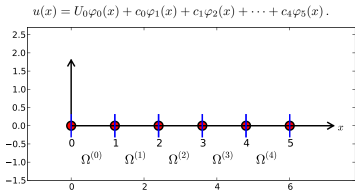
It is easy to verify that $B(x_i) = \sum_{j \in I_b} U_j \varphi_j(x_i) = U_i$. The unknown function can then be written as

$$u(x) = \sum_{j \in I_b} U_j \varphi_j(x) + \sum_{j \in \mathcal{I}_s} c_j \varphi_{\nu(j)},$$

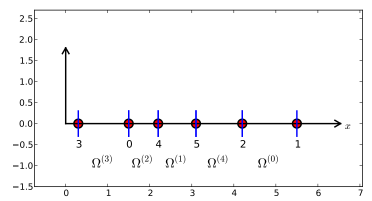
where $\nu(j)$ maps unknown number j in the equation system to node ν can easily show that with this u, a Dirichlet condition $u(x_k) = U_k$ is fu

$$u(x_k) = \sum_{j \in I_b} U_j \qquad \underbrace{\varphi_j(x)}_{\neq 0 \text{ only for } j = k} + \sum_{j \in \mathcal{I}_s} c_j \underbrace{\varphi_{\nu(j)}(x_k)}_{=0, \ k \notin \mathcal{I}_s} = U_k$$

Some examples will further clarify the notation. With a regular left-numbering of nodes in a mesh with P1 elements, and Dirichlet condi x=0, we use finite element basis functions associated with the nodes 1, 2, implying that $\nu(j)=j+1, j=0,\ldots,N$, where $N=N_n-1$. For the pamesh below the expansion becomes



Here is a mesh with an irregular cell and node numbering:



Say we in this latter mesh have Dirichlet conditions on the left-most and ght-most node, with numbers 3 and 1, respectively. Then we can number the nknowns at the interior nodes from left to right, giving $\nu(0) = 0$, $\nu(1) = 4$, (2) = 5, $\nu(3) = 2$. This gives

$$B(x) = U_3\varphi_3(x) + U_1\varphi_1(x),$$

 nd

$$u(x) = B(x) + \sum_{j=0}^{3} c_j \varphi_{\nu(j)} = U_3 \varphi_3 + U_1 \varphi_1 + c_0 \varphi_0 + c_1 \varphi_4 + c_2 \varphi_5 + c_3 \varphi_2.$$

Switching to the more standard case of left-to-right numbering and boundary onditions u(0) = C, u(L) = D, we have $N = N_n - 2$ and

$$u(x) = C\varphi_0 + D\varphi_{N_n} + \sum_{j \in \mathcal{I}_s} c_j \varphi_{j+1}$$

= $C\varphi_0 + D\varphi_{N_n} + c_0 \varphi_1 + c_1 \varphi_2 + \dots + c_N \varphi_{N_n-1}$.

The idea of constructing B described here generalizes almost trivially to 2D nd 3D problems: $B = \sum_{j \in I_b} U_j \varphi_j$, where I_b is the index set containing the umbers of all the nodes on the boundaries where Dirichlet values are prescribed.

4.2 Example on computing with finite element-based a boundary function

et us see how the model problem -u'' = 2, u(0) = C, u(L) = D, is affected by B(x) to incorporate boundary values. Inserting the expression

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x)$$

 $1 - (u'', \psi_i) = (f, \psi_i)$ and integrating by parts results in a linear system with

$$A_{i,j} = \int_0^L \psi_i'(x)\psi_j'(x) dx, \quad b_i = \int_0^L (f(x) - B'(x))\psi_i(x) dx.$$

We choose $\psi_i = \varphi_{i+1}$, $i = 0, ..., N = N_n - 2$ if the node numbering is f to right. (Later we also need the assumption that the cells too are nu from left to right.) The boundary function becomes

$$B(x) = C\varphi_0(x) + D\varphi_{N_n}(x).$$

The expansion for u(x) is

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \varphi_{j+1}(x).$$

We can write the matrix and right-hand side entries as

$$A_{i-1,j-1} = \int_0^L \varphi_i'(x)\varphi_j'(x) \, \mathrm{d}x, \quad b_{i-1} = \int_0^L (f(x) - C\varphi_0'(x) - D\varphi_{N_n}'(x))\varphi$$

for $i, j = 1, ..., N+1 = N_n - 1$. Note that we have here used $B' = C\varphi'_0$

Computations in physical coordinates. Most of the terms in th system have already been computed so we concentrate on the new contraction from the boundary function. The integral $C\int_0^L \varphi_0'(x))\varphi_i(x)\,\mathrm{d}x$ can on nonzero contribution from the first cell, $\Omega^{(0)}=[x_0,x_1]$ since $\varphi_0'(x)=$ other cells. Moreover, $\varphi_0'(x)\varphi_i(x)\,\mathrm{d}x\neq 0$ only for i=0 and i=1 (but excluded), since $\varphi_i=0$ on the first cell if i>1. With a similar reaso realize that $D\int_0^L \varphi_{N_n}'(x))\varphi_i(x)\,\mathrm{d}x$ can only get a nonzero contribution f last cell. From the explanations of the calculations in Section 3.6 we that

$$\int_0^L \varphi_0'(x)\varphi_1(x) \, \mathrm{d}x = \frac{1}{h} \cdot \frac{1}{h} = -\frac{1}{2}, \quad \int_0^L \varphi_{N_n}'(x)\varphi_{N_n-1}(x) \, \mathrm{d}x = \frac{1}{h} \cdot \frac{1}{h}$$

The extra boundary term because of B(x) boils down to adding C/2 to -D/2 to b_N .

Cellwise computations on the reference element. As an equivalent ternative, we now turn to cellwise computations. The element matrix vectors are calculated as Section 13.4, so we concentrate on the impact new term involving B(x). We observe that $C\varphi'_0 = 0$ on all cells except and $D\varphi'_{N_n} = 0$ on all cells except $e = N_e$. In this case there is only one u in these cells since u(0) and u(L) are prescribed, so the element vector one entry. The entry for the last cell, $e = N_e$, becomes

$$\tilde{b}_{0}^{(e)} = \int_{-1}^{1} \left(f - D \frac{2}{h} \frac{d\tilde{\varphi}_{1}}{dX} \right) \tilde{\varphi}_{0} \frac{h}{2} dX = \left(\frac{h}{2} (2 - D \frac{2}{h} \frac{1}{2}) \int_{-1}^{1} \tilde{\varphi}_{0} dX = h - D/2 \right).$$

imilar computations on the first cell yield

$$\tilde{b}_0^{(0)} = \int_{-1}^1 \left(f - C \frac{2}{h} \frac{d\tilde{\varphi}_0}{dX} \right) \tilde{\varphi}_1 \frac{h}{2} dX = \left(\frac{h}{2} (2 + C \frac{2}{h} \frac{1}{2}) \int_{-1}^1 \tilde{\varphi}_1 dX = h + C/2 \right).$$

/hen assembling these contributions, we see that b_0 gets right-hand side of the near system gets an extra term C/2 and b_N gets -D/2, as in the computations 1 the physical domain.

4.3 Modification of the linear system

rom an implementational point of view, there is a convenient alternative to dding the B(x) function and using only the basis functions associated with odes where u is truly unknown. Instead of seeking

$$u(x) = \sum_{j \in I_b} U_j \varphi_j(x) + \sum_{j \in \mathcal{I}_s} c_j \varphi_{\nu(j)}(x), \tag{181}$$

e use the sum over all degrees of freedom, including the known boundary values:

$$u(x) = \sum_{j \in \mathcal{I}_s} c_j \varphi_j(x). \tag{182}$$

ote that the collections of unknowns $\{c_i\}_{i\in\mathcal{I}_s}$ in (181) and (182) are different: ι (181) N counts the number of nodes where u is not known, while in (181) N punts all the nodes $(N=N_n)$.

The idea is to compute the entries in the linear system as if no Dirichlet alues are prescribed. Afterwards, we modify the linear system to ensure that ne known c_i values are incorporated.

A potential problem arises for the boundary term $[u'v]_0^L$ from the integration y parts: imagining no Dirichlet conditions means that we no longer require =0 at Dirichlet points, and the boundary term is then nonzero at these oints. However, when we modify the linear system, we will erase whatever the antribution from $[u'v]_0^L$ should be at the Dirichlet points in the right-hand side \mathbb{I} the linear system. We can therefore safely forget $[u'v]_0^L$ at any point where a virichlet condition applies.

Computations in the physical system. Let us redo the computations in the example in Section 14.1. We solve -u'' = 2 with u(0) = 0 and u(L) = D. the expressions for $A_{i,j}$ and b_i are the same, but the numbering is different as the numbering of unknowns and nodes now coincide:

$$A_{i,j} = \int_0^L \varphi_i'(x)\varphi_j'(x) \, \mathrm{d}x, \quad b_i = \int_0^L f(x)\varphi_i(x) \, \mathrm{d}x,$$

for $i, j = 0, ..., N = N_n$. The integrals involving basis functions correspond interior mesh nodes, $i, j = 1, ..., N_n - 1$, are obviously the same as We concentrate on the contributions from φ_0 and φ_{N_n} :

$$A_{0,0} = \int_0^L (\varphi_0')^2 dx = \int_0^{x_1} = (\varphi_0')^2 dx \frac{1}{h},$$

$$A_{0,1} = \int_0^L \varphi_0' \varphi_1' dx = \int_0^{x_1} \varphi_0' \varphi_1' dx = -\frac{1}{h},$$

$$A_{N,N} = \int_0^L (\varphi_0')^2 dx = \int_{x_{N_n-1}}^{x_{N_n}} (\varphi_0')^2 dx = \frac{1}{h},$$

$$A_{N,N-1} = \int_0^L (\varphi_0')^2 dx = \int_{x_{N_n-1}}^{x_{N_n}} (\varphi_0')^2 dx = -\frac{1}{h}.$$

The new terms on the right-hand side are also those involving φ_0 as

$$b_0 = \int_0^L 2\varphi_0(x) \, dx = \int_0^{x_1} 2\varphi_0(x) \, dx = h,$$

$$b_N = \int_0^L 2\varphi_{N_n} \, dx = \int_{x_{N_n}}^{x_{N_n}} 2\varphi_{N_n} \, dx = h.$$

The complete matrix system, involving all degrees of freedom, takes t

$$\begin{pmatrix}
1 & -1 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\
-1 & 2 & -1 & \ddots & & & & \vdots \\
0 & -1 & 2 & -1 & \ddots & & & \vdots \\
\vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
\vdots & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & -1 \\
0 & \cdots & \cdots & \cdots & \cdots & 0 & -1 & 1
\end{pmatrix}
\begin{pmatrix}
c_0 \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N
\end{pmatrix} = \begin{pmatrix}
\vdots \\ c_N
\end{pmatrix}$$

Incorporation of Dirichlet values can now be done by replacing the f last equation by $c_0 = 0$ and $c_N = D$. This action changes the system to

$$\begin{pmatrix}
h & 0 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\
-1 & 2 & -1 & \ddots & & & & \vdots \\
0 & -1 & 2 & -1 & \ddots & & & \vdots \\
\vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
\vdots & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & & \ddots & \ddots & \ddots & \ddots & -1 \\
0 & \cdots & \cdots & \cdots & \cdots & 0 & 0 & 0 & h
\end{pmatrix}
\begin{pmatrix}
c_0 \\ \vdots \\ c_N
\end{pmatrix} = \begin{pmatrix}
0 \\ 2h \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ 2h \\ D
\end{pmatrix}$$
(184)

Note that because we do not require $\varphi_i(0) = 0$ and $\varphi_i(L)$, $i \in \mathcal{I}_s$, the boundary $\operatorname{rm} [u'v]_0^L$ gives in principle contributions $u'(0)\varphi_0(0)$ to b_0 and $u'(L)\varphi_N(L)$ to $v(u'\varphi_i)$ vanishes for x=0 or x=L for $i=1,\ldots,N-1$). Nevertheless, we asse these contributions in b_0 and b_N and insert boundary values instead. This regument shows why we can drop computing $[u'v]_0^L$ at Dirichlet nodes when we nplement the Dirichlet values by modifying the linear system.

4.4 Symmetric modification of the linear system

he original matrix system (172) is symmetric, but the modifications in (184) estroy the symmetry. Our described modification will in general destroy an itial symmetry in the matrix system. This is not a particular computational isadvantage for tridiagonal systems arising in 1D problems, but may be more grious in 2D and 3D problems when the systems are large and exploiting matrix can be important for halving the storage demands, speeding up omputations, and/or making the solution algorithm more robust. Therefore, a alternative modification which preserves symmetry is frequently applied.

Let c_k be a coefficient corresponding to a known value $u(x_k) = U_k$. We ant to replace equation k in the system by $c_k = U_k$, i.e., insert zeroes in row umber k in the coefficient matrix, set 1 on the diagonal, and replace b_k by U_k . symmetry-preserving modification consists in first subtracting column number in the coefficient matrix, i.e., $A_{i,k}$ for $i \in \mathcal{I}_s$, times the boundary value U_k , om the right-hand side: $b_i \leftarrow b_i - A_{i,k}U_k$. Then we put zeroes in row number and column number k in the coefficient matrix, and finally set $b_k = U_k$. The teps in algorithmic form becomes

1.
$$b_i \leftarrow b_i - A_{ik}U_k$$
 for $i \in \mathcal{I}_s$

2.
$$A_{i,k} = A_{k,i} = 0$$
 for $i \in \mathcal{I}_s$

3.
$$A_{k,k} = 1$$

4.
$$b_i = U_k$$

This modification goes as follows for the specific linear system writter (183) in Section 14.3. First we subtract the first column in the coefficient times the boundary value, from the right-hand side. Because $c_0 =$ subtraction has no effect. Then we subtract the last column, times the boundary D, from the right-hand side. This action results in $b_{N-1} = 2h$ and $b_N = h - 2D/h$. Thereafter, we place zeros in the first and last 1 column in the coefficient matrix and 1 on the two corresponding diagonal Finally, we set $b_0 = 0$ and $b_N = D$. The result becomes

$$\begin{pmatrix}
h & 0 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\
0 & 2 & -1 & \ddots & & & & & \vdots \\
0 & -1 & 2 & -1 & \ddots & & & & \vdots \\
\vdots & \ddots & \ddots & \ddots & \ddots & \ddots & & \vdots \\
\vdots & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
0 & \cdots & \cdots & \cdots & \cdots & \cdots & 0 & 0 & h
\end{pmatrix}
\begin{pmatrix}
c_0 \\ \vdots \\ c_N
\end{pmatrix} = \begin{pmatrix}
0 \\ 2h \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ 2h + D
\end{pmatrix}$$

14.5 Modification of the element matrix and vector

The modifications of the global linear system can alternatively be done element matrix and vector. (The assembled system will get the value r main diagonal if n elements contribute to the same unknown, but the swill also appear on the right-hand side and hence cancel out.)

We have, in the present computational example, the element mat vector (177). The modifications are needed in cells where one of the of freedom is known. Here, this means the first and last cell. We compelement matrix and vector as there are no Dirichlet conditions. The betterm $[u'v]_0^L$ is simply forgotten at nodes that have Dirichlet conditions the modification of the element vector will anyway erase the contribution the boundary term. In the first cell, local degree of freedom number 0 is and the modification becomes

$$\tilde{A}^{(0)} = A = \frac{1}{h} \begin{pmatrix} h & 0 \\ -1 & 1 \end{pmatrix}, \quad \tilde{b}^{(0)} = \begin{pmatrix} 0 \\ h \end{pmatrix}.$$

In the last cell we set

$$\tilde{A}^{(N_e)} = A = \frac{1}{h} \begin{pmatrix} 1 & -1 \\ 0 & h \end{pmatrix}, \quad \tilde{b}^{(N_e)} = \begin{pmatrix} h \\ D \end{pmatrix}. \tag{187}$$

We can also perform the symmetric modification. This operation affects only ne last cell with a nonzero Dirichlet condition. The algorithm is the same as or the global linear system, resulting in

$$\tilde{A}^{(N-1)} = A = \frac{1}{h} \begin{pmatrix} h & 0 \\ 0 & 1 \end{pmatrix}, \quad \tilde{b}^{(N-1)} = \begin{pmatrix} h + D/h \\ D \end{pmatrix}. \tag{188}$$

he reader is encouraged to assemble the element matrices and vectors and check nat the result coincides with the system (185).

5 Boundary conditions: specified derivative

uppose our model problem -u''(x) = f(x) features the boundary conditions '(0) = C and u(L) = D. As already indicated in Section 12, the former ondition can be incorporated through the boundary term that arises from itegration by parts. This details of this method will now be illustrated in the ontext of finite element basis functions.

5.1 The variational formulation

tarting with the Galerkin method,

$$\int_0^L (u''(x) + f(x))\psi_i(x) dx = 0, \quad i \in \mathcal{I}_s,$$

itegrating $u''\psi_i$ by parts results in

$$\int_0^L u'(x)' \psi_i'(x) \, \mathrm{d}x - (u'(L)\psi_i(L) - u'(0)\psi_i(0)) = \int_0^L f(x)\psi_i(x) \, \mathrm{d}x, \quad i \in \mathcal{I}_s.$$

The first boundary term, $u'(L)\psi_i(L)$, vanishes because u(L) = D. There are vo arguments for this result, explained in detail below. The second boundary erm, $u'(0)\psi_i(0)$, can be used to implement the condition u'(0) = C, provided $i(0) \neq 0$ for some i (but with finite elements we fortunately have $\psi_0(0) = 1$). he variational form of the differential equation then becomes

$$\int_0^L u'(x)\varphi_i'(x) dx + C\varphi_i(0) = \int_0^L f(x)\varphi_i(x) dx, \quad i \in \mathcal{I}_s.$$

15.2 Boundary term vanishes because of the test fun-

At points where u is known we may require ψ_i to vanish. Here, u(L) = then $\psi_i(L) = 0$, $i \in \mathcal{I}_s$. Obviously, the boundary term $u'(L)\psi_i(L)$ then

The set of basis functions $\{\psi_i\}_{i\in\mathcal{I}_s}$ contains in this case all the finite basis functions on the mesh, expect the one that is 1 at x=L. The function that is left out is used in a boundary function B(x) instead. left-to-right numbering, $\psi_i = \varphi_i$, $i = 0, ..., N_n - 1$, and $B(x) = D\varphi_{N_n}$:

$$u(x) = D\varphi_{N_n}(x) + \sum_{i=0}^{N=N_n-1} c_j \varphi_j(x).$$

Inserting this expansion for u in the variational form (15.1) leads to the system

$$\sum_{j=0}^{N} \left(\int_{0}^{L} \varphi_i'(x) \varphi_j'(x) \, \mathrm{d}x \right) c_j = \int_{0}^{L} \left(f(x) \varphi_i(x) - D \varphi_{N_n}'(x) \varphi_i(x) \right) \, \mathrm{d}x - C$$

for
$$i = 0, ..., N = N_n - 1$$
.

15.3 Boundary term vanishes because of linear symmodifications

We may, as an alternative to the approach in the previous section, use $\{\psi_i\}_{i\in\mathcal{I}_s}$ which contains all the finite element functions on the mesh: $i=0,\ldots,N_n=N$. In this case, $u'(L)\psi_i(L)=u'(L)\varphi_i(L)\neq 0$ for corresponding to the boundary node at x=L (where $\varphi_i=1$). The nuthis node is $i=N_n=N$ if a left-to-right numbering of nodes is utilized

However, even though $u'(L)\varphi_N(L) \neq 0$, we do not need to compute the For i < N we realize that $\varphi_i(L) = 0$. The only nonzero contribution right-hand side from the affects b_N (i = N). Without a boundary function must implement the condition u(L) = D by the equivalent statement and modify the linear system accordingly. This modification will erase row and replace b_N by another value. Any attempt to compute the botterm $u'(L)\varphi_N(L)$ and store it in b_N will be lost. Therefore, we can safel about boundary terms corresponding to Dirichlet boundary conditions alwe use the methods from Section 14.3 or Section 14.4.

The expansion for u reads

$$u(x) = \sum_{j \in \mathcal{I}_s} c_j \varphi_j(x), \quad B(x) = D\varphi_N(x),$$

with $N = N_n$. Insertion in the variational form (15.1) leads to the linear

$$\sum_{j \in \mathcal{I}_s} \left(\int_0^L \varphi_i'(x) \varphi_j'(x) \, \mathrm{d}x \right) c_j = \int_0^L \left(f(x) \varphi_i(x) \right) \, \mathrm{d}x - C \varphi_i(0), \quad i \in \mathcal{I}_s. \tag{190}$$

fter having computed the system, we replace the last row by $c_N=D$, either raightforwardly as in Section refreffem:deq:1D:fem:essBC:Bfunc:modsys or in a /mmetric fashion as in Section refreffem:deq:1D:fem:essBC:Bfunc:modsys:symm. hese modifications can also be performed in the element matrix and vector for ne right-most cell.

5.4 Direct computation of the global linear system

/e now turn to actual computations with P1 finite elements. The focus is on ow the linear system and the element matrices and vectors are modified by the ondition u'(0) = C.

Consider first the approach where Dirichlet conditions are incorporated by a C(x) function and the known degree of freedom C_{N_n} is left out from the linear stem (see Section 15.2). The relevant formula for the linear system is given y (189). There are three differences compared to the extensively computed ase where u(0)=0 in Sections 13.2 and 13.4. First, because we do not have a ririchlet condition at the left boundary, we need to extend the linear system (172) ith an equation associated with the node $x_0=0$. According to Section 14.3, this stension consists of including $A_{0,0}=1/h$, $A_{0,1}=-1/h$, and $b_0=h$. For i>0 e have $A_{i,i}=2/h$, $A_{i-1,i}=A_{i,i+1}=-1/h$. Second, we need to include the stra term $-C\varphi_i(0)$ on the right-hand side. Since all $\varphi_i(0)=0$ for $i=1,\ldots,N$, is term reduces to $-C\varphi_0(0)=-C$ and affects only the first equation (i=0). We simply add -C to b_0 such that $b_0=h-C$. Third, the boundary term $\int_0^L D\varphi_{N_n}(x)\varphi_i\,\mathrm{d}x$ must be computed. Since $i=0,\ldots,N=N_n-1$, this itegral can only get a nonzero contribution with $i=N_n-1$ over the last cell. he result becomes -Dh/6. The resulting linear system can be summarized in ne form

$$\begin{pmatrix}
1 & -1 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\
-1 & 2 & -1 & \ddots & & & & \vdots \\
0 & -1 & 2 & -1 & \ddots & & & \vdots \\
\vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
\vdots & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & -1 \\
0 & \cdots & \cdots & \cdots & \cdots & 0 & -1 & 2
\end{pmatrix}
\begin{pmatrix}
c_0 \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N
\end{pmatrix} = \begin{pmatrix}
c_0 \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N
\end{pmatrix}$$

Next we consider the technique where we modify the linear system to porate Dirichlet conditions (see Section 15.3). Now $N=N_n$. The two differom the case above is that the $-\int_0^L D\varphi_{N_n}\varphi_i\,\mathrm{d}x$ term is left out of the right side and an extra last row associated with the node $x_{N_n}=L$ where the I condition applies is appended to the system. This last row is anyway result by the condition $C_N=D$ or this condition can be incorporated in a synfashion. Using the simplest, former approach gives

$$\begin{pmatrix}
1 & -1 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\
-1 & 2 & -1 & \ddots & & & & \vdots \\
0 & -1 & 2 & -1 & \ddots & & & \vdots \\
\vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\
\vdots & & 0 & -1 & 2 & -1 & \ddots & \vdots \\
\vdots & & & \ddots & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\
\vdots & & & & \ddots & \ddots & \ddots & \ddots & 0
\end{bmatrix} = \begin{pmatrix} h - 2 \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N \end{pmatrix}$$

15.5 Cellwise computations

Now we compute with one element at a time, working in the reference consystem $X \in [-1,1]$. We need to see how the u'(0) = C condition the element matrix and vector. The extra term $-C\varphi_i(0)$ in the var formulation only affects the element vector in the first cell. On the refere $-C\varphi_i(0)$ is transformed to $-C\tilde{\varphi}_r(-1)$, where r counts local degrees of

We have $\tilde{\varphi}_0(-1) = 1$ and $\tilde{\varphi}_1(-1) = 0$ so we are left with the contribution $C\tilde{\varphi}_0(-1) = -C$ to $\tilde{b}_0^{(0)}$:

$$\tilde{A}^{(0)} = A = \frac{1}{h} \begin{pmatrix} 1 & 1 \\ -1 & 1 \end{pmatrix}, \quad \tilde{b}^{(0)} = \begin{pmatrix} h - C \\ h \end{pmatrix}.$$
 (193)

o other element matrices or vectors are affected by the $-C\varphi_i(0)$ boundary erm.

There are two alternative ways of incorporating the Dirichlet condition. ollowing Section 15.2, we get a 1×1 element matrix in the last cell and an ement vector with an extra term containing D:

$$\tilde{A}^{(e)} = \frac{1}{h} \begin{pmatrix} 1 \end{pmatrix}, \quad \tilde{b}^{(e)} = h \begin{pmatrix} 1 - D/6 \end{pmatrix},$$
 (194)

Alternatively, we include the degree of freedom at the node with u specified he element matrix and vector must then be modified to constrain the $\tilde{c}_1 = c_N$ alue at local node r = 1:

$$\tilde{A}^{(N_e)} = A = \frac{1}{h} \begin{pmatrix} 1 & 1 \\ 0 & 1 \end{pmatrix}, \quad \tilde{b}^{(N_e)} = \begin{pmatrix} h \\ D \end{pmatrix}.$$
 (195)

6 Implementation

is tempting to create a program with symbolic calculations to perform all the seps in the computational machinery, both for automating the work and for ocumenting the complete algorithms. As we have seen, there are quite many etails involved with finite element computations and incorporation of boundary onditions. An implementation will also act as a structured summary of all these etails.

6.1 Global basis functions

/e first consider implementations when ψ_i are global functions are hence different om zero on most of $\Omega=[0,L]$ so all integrals need integration over the entire omain. Since the expressions for the entries in the linear system depend on the ifferential equation problem being solved, the user must supply the necessary ormulas via Python functions. The implementations here attempt to perform /mbolic calculations, but fall back on numerical computations if the symbolic nes fail.

The user must prepare a function integrand_lhs(psi, i, j) for returning ne integrand of the integral that contributes to matrix entry (i, j). The psi wiable is a Python dictionary holding the basis functions and their derivatives symbolic form. More precisely, psi[q] is a list of

$$\left\{\frac{d^q\psi_0}{dx^q},\dots,\frac{d^q\psi_N}{dx^q}\right\}.$$

Similarly, integrand_rhs(psi, i) returns the integrand for entry num the right-hand side vector.

Since we also have contributions to the right-hand side vector, and pot also the matrix, from boundary terms without any integral, we introd additional functions, boundary_lhs(psi, i, j) and boundary_rhs(] for returning terms in the variational formulation that are not to be int over the domain Ω . Examples shown later will explain in more detail he user-supplied function may look like.

The linear system can be computed and solved symbolically by the function:

```
import sympy as sp
def solve(integrand_lhs, integrand_rhs, psi, Omega,
          boundary_lhs=None, boundary_rhs=None):
    N = len(psi[0]) - 1
    A = sp.zeros((N+1, N+1))
    b = sp.zeros((N+1, 1))
    x = sp.Symbol('x')
    for i in range(N+1):
        for j in range(i, N+1):
            integrand = integrand_lhs(psi, i, j)
            I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
            if boundary lhs is not None:
                I += boundary_lhs(psi, i, j)
            A[i,j] = A[j,i] = I # assume symmetry
        integrand = integrand_rhs(psi, i)
        I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
        if boundary rhs is not None:
            I += boundary_rhs(psi, i)
        b[i,0] = I
    c = A.LUsolve(b)
    u = sum(c[i,0]*psi[0][i]  for i in range(len(psi[0])))
```

Not surprisingly, symbolic solution of differential equations, discretiz Galerkin or least squares method with global basis functions, is of limited beyond the simplest problems, because symbolic integration might be veconsuming or impossible, not only in sympy but also in WolframAlpha²² applies the perhaps most powerful symbolic integration software available Mathematica). Numerical integration as an option is therefore desirable

The extended solve function below tries to combine symbolic and m integration. The latter can be enforced by the user, or it can be invoked non-successful symbolic integration (being detected by an Integral of the result of the integration in sympy). Note that for a numerical integrable symbolic expressions must be converted to Python functions (using lam and the expressions cannot contain other symbols than x. The real solve in the varform1D.py²³ file has error checking and meaningful error mes such cases. The solve code below is a condensed version of the real or

²²http://wolframalpha.com

²³http://tinyurl.com/nm5587k/fem/varform1D.py

ne purpose of showing how to automate the Galerkin or least squares method or solving differential equations in 1D with global basis functions:

```
lef solve(integrand lhs, integrand rhs, psi, Omega,
         boundary_lhs=None, boundary_rhs=None, symbolic=True):
   N = len(psi[0]) - 1
   A = sp.zeros((N+1, N+1))
   b = sp.zeros((N+1, 1))
   x = sp.Symbol('x')
   for i in range(N+1):
       for j in range(i, N+1):
           integrand = integrand_lhs(psi, i, j)
           if symbolic:
               I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
               if isinstance(I, sp.Integral):
                   symbolic = False # force num.int. hereafter
           if not symbolic:
               integrand = sp.lambdify([x], integrand)
               I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]])
           if boundary lhs is not None:
               I += boundary_lhs(psi, i, j)
           A[i,j] = A[j,i] = I
       integrand = integrand rhs(psi, i)
       if symbolic:
           I = sp.integrate(integrand, (x, Omega[0], Omega[1]))
           if isinstance(I, sp.Integral):
               symbolic = False
       if not symbolic:
           integrand = sp.lambdify([x], integrand)
           I = sp.mpmath.quad(integrand, [Omega[0], Omega[1]])
       if boundary_rhs is not None:
           I += boundary rhs(psi, i)
       b[i.0] = I
   c = A.LUsolve(b)
   u = sum(c[i,0]*psi[0][i]  for i in range(len(psi[0])))
   return u
```

6.2 Example: constant right-hand side

o demonstrate the code above, we address

$$-u''(x) = b$$
, $x \in \Omega = [0, 1]$, $u(0) = 1$, $u(1) = 0$,

ith b as a (symbolic) constant. A possible basis for the space V is $\psi_i(x) = i^{+1}(1-x)$, $i \in \mathcal{I}_s$. Note that $\psi_i(0) = \psi_i(1) = 0$ as required by the Dirichlet onditions. We need a B(x) function to take care of the known boundary values f(u). Any function $B(x) = 1 - x^p$, $p \in \mathbb{R}$, is a candidate, and one arbitrary noise from this family is $B(x) = 1 - x^3$. The unknown function is then written

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_0} c_j \psi_j(x).$$

Let us use the Galerkin method to derive the variational formulation. Multilying the differential equation by v and integrate by parts yield

$$\int_0^1 u'v' \, \mathrm{d}x = \int_0^1 fv \, \mathrm{d}x \quad \forall v \in V,$$

and with $u = B + \sum_{j} c_{j} \psi_{j}$ we get the linear system

$$\sum_{j \in \mathcal{I}_s} \left(\int_0^1 \psi_i' \psi_j' \, \mathrm{d}x \right) c_j = \int_0^1 (f - B') \psi_i \, \mathrm{d}x, \quad i \in \mathcal{I}_s.$$

The application can be coded as follows in sympy:

```
x, b = sp.symbols('x b')
f = b
B = 1 - x**3
dBdx = sp.diff(B, x)
# Compute basis functions and their derivatives
N = 3
psi = \{0: [x**(i+1)*(1-x) \text{ for i in } range(N+1)]\}
psi[1] = [sp.diff(psi_i, x) for psi_i in psi[0]]
def integrand_lhs(psi, i, j):
    return psi[1][i]*psi[1][j]
def integrand rhs(psi, i):
    return f*psi[0][i] - dBdx*psi[1][i]
Omega = [0, 1]
u_bar = solve(integrand_lhs, integrand_rhs, psi, Omega,
              verbose=True, symbolic=True)
u = B + u bar
print 'solution u:', sp.simplify(sp.expand(u))
```

The printout of u reads -b*x**2/2 + b*x/2 - x + 1. Note that expa and then simplifying is in the present case necessary to get a comparexpression with sympy. A non-expanded u might be preferable in othe this depends on the problem in question.

The exact solution $u_{\rm e}(x)$ can be derived by some sympy code that follows the examples in Section 11.2. The idea is to integrate -u''=b to determine the integration constants from the boundary conditions:

```
C1, C2 = sp.symbols('C1 C2')  # integration constants
f1 = sp.integrate(f, x) + C1
f2 = sp.integrate(f1, x) + C2
# Find C1 and C2 from the boundary conditions u(0)=0, u(1)=1
s = sp.solve([u_e.subs(x,0) - 1, u_e.subs(x,1) - 0], [C1, C2])
# Form the exact solution
u_e = -f2 + s[C1]*x + s[C2]
print 'analytical solution:', u_e
print 'error:', sp.simplify(sp.expand(u - u_e))
```

The last line prints 0, which is not surprising when $u_{\rm e}(x)$ is a parabola approximate u contains polynomials up to degree 4. It suffices to have i.e., polynomials of degree 2, to recover the exact solution.

We can play around with the code and test that with $f \sim x^p$, the solution is polynomial of degree p+2, and N=p+1 guarantees that the approximate plution is exact.

Although the symbolic code is capable of integrating many choices of f(x), ne symbolic expressions for u quickly become lengthy and non-informative, numerical integration in the code, and hence numerical answers, have the reatest application potential.

6.3 Finite elements

nplementation of the finite element algorithms for differential equations follows closely the algorithm for approximation of functions. The new additional agredients are

- 1. other types of integrands (as implied by the variational formulation)
- 2. additional boundary terms in the variational formulation for Neumann boundary conditions
- modification of element matrices and vectors due to Dirichlet boundary conditions

oint 1 and 2 can be taken care of by letting the user supply functions defining ne integrands and boundary terms on the left- and right-hand side of the quation system:

```
integrand_lhs(phi, r, s, x)
poundary_lhs(phi, r, s, x)
integrand_rhs(phi, r, x)
poundary_rhs(phi, r, x)
```

ere, phi is a dictionary where phi[q] holds a list of the derivatives of order of the basis functions at the an evaluation point; \mathbf{r} and \mathbf{s} are indices for the presponding entries in the element matrix and vector, and \mathbf{x} is the global pordinate value corresponding to the current evaluation point.

Given a mesh represented by vertices, cells, and dof_map as explained bere, we can write a pseudo Python code to list all the steps in the computational lgorithm for finite element solution of a differential equation.

```
CDeclare global matrix and rhs: A, b>
ior e in range(len(cells)):

# Compute element matrix and vector
n = len(dof_map[e]) # no of dofs in this element
h = vertices[cells[e][1]] - vertices[cells[e][1]]
<Declare element matrix and vector: A_e, b_e>

# Integrate over the reference cell
points, weights = <numerical integration rule>
for X, w in zip(points, weights):
```

```
phi = <basis functions and derivatives at X>
       detJ = h/2
       x = <affine mapping from X>
       for r in range(n):
           for s in range(n):
               A_e[r,s] + integrand_lhs(phi, r, s, x)*detJ*w
           b e[r] += integrand rhs(phi, r, x)*detJ*w
    # Add boundary terms
   for r in range(n):
       for s in range(n):
            A_e[r,s] += boundary_lhs(phi, r, s, x)*detJ*w
       b e[r] += boundary rhs(phi, r, x)*detJ*w
    # Incorporate essential boundary conditions
   for r in range(n):
       global_dof = dof_map[e][r]
        if global dof in essbc dofs:
            # dof r is subject to an essential condition
            value = essbc docs[global dof]
            # Symmetric modification
           b_e -= value*A_e[:,r]
            A = [r, :] = 0
           A_e[:,r] = 0
           A_e[r,r] = 1
           b_e[r] = value
    # Assemble
   for r in range(n):
       for s in range(n):
           A[dof_map[e][r], dof_map[e][r]] += A_e[r,s]
       b[dof map[e][r] += b e[r]
<solve linear system>
```

17 Variational formulations in 2D and 3D

The major difference between deriving variational formulations in 2D compared to 1D is the rule for integrating by parts. A typical second-ord in a PDE may be written in dimension-independent notation as

$$\nabla^2 u$$
 or $\nabla \cdot (a(\boldsymbol{x})\nabla u)$.

The explicit forms in a 2D problem become

$$\nabla^2 u = \nabla \cdot \nabla u = \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2},$$

and

$$\nabla \cdot (a(\boldsymbol{x})\nabla u) = \frac{\partial}{\partial x} \left(a(x,y) \frac{\partial u}{\partial x} \right) + \frac{\partial}{\partial y} \left(a(x,y) \frac{\partial u}{\partial y} \right) .$$

We shall continue with the latter operator as the form arises from just a=1.

The general rule for integrating by parts is often referred to as Green's first lentity 24 :

$$-\int_{\Omega} \nabla \cdot (a(\boldsymbol{x})\nabla u)v \, d\boldsymbol{x} = \int_{\Omega} a(\boldsymbol{x})\nabla u \cdot \nabla v \, d\boldsymbol{x} - \int_{\partial\Omega} a \frac{\partial u}{\partial n} v \, ds, \qquad (197)$$

here $\partial\Omega$ is the boundary of Ω and $\partial u/\partial n = n \cdot \nabla u$ is the derivative of u in the utward normal direction, n being an outward unit normal to $\partial\Omega$. The integrals $\alpha_{\Omega}(x)$ are area integrals in 2D and volume integrals in 3D, while $\alpha_{\Omega}(x)$ ds is a ne integral in 2D and a surface integral in 3D.

Let us divide the boundary into two parts:

- $\partial \Omega_N$, where we have Neumann conditions $-a \frac{\partial u}{\partial n} = g$, and
- $\partial \Omega_D$, where we have Dirichlet conditions $u = u_0$.

he test functions v are required to vanish on $\partial \Omega_D$.

xample. Here is a quite general, stationary, linear PDE arising in many roblems:

$$\boldsymbol{v} \cdot \nabla u + \alpha u = \nabla \cdot (a\nabla u) + f, \quad \boldsymbol{x} \in \Omega,$$
 (198)

$$u = u_0, \quad \boldsymbol{x} \in \partial \Omega_D,$$
 (199)

$$-a\frac{\partial u}{\partial n} = g, \quad \boldsymbol{x} \in \partial\Omega_N. \tag{200}$$

he vector field v and the scalar functions a, α , f, u_0 , and g may vary with the patial coordinate x and must be known.

Such a second-order PDE needs exactly one boundary condition at each point f the boundary, so $\partial \Omega_N \cup \partial \Omega_D$ must be the complete boundary $\partial \Omega$.

Assume that the boundary function $u_0(x)$ is defined for all $x \in \Omega$. The nknown function can then be expanded as

$$u = B + \sum_{j \in \mathcal{I}_s} c_j \psi_j, \quad B = u_0.$$

he variational formula is obtained from Galerkin's method, which technically nplies multiplying the PDE by a test function v and integrating over Ω :

$$\int_{\Omega} (\boldsymbol{v} \cdot \nabla u + \alpha u) v \, \mathrm{d}x = \int_{\Omega} \nabla \cdot (a \nabla u) \, \mathrm{d}x + \int_{\Omega} f v \, \mathrm{d}x.$$

he second-order term is integrated by parts, according to

$$\int_{\Omega} \nabla \cdot (a \nabla u) v \, dx = -\int_{\Omega} a \nabla u \cdot \nabla v \, dx + \int_{\partial \Omega} a \frac{\partial u}{\partial n} v \, ds.$$

The variational form now reads

$$\int_{\Omega} (\boldsymbol{v} \cdot \nabla u + \alpha u) v \, \mathrm{d}x = -\int_{\Omega} a \nabla u \cdot \nabla v \, \mathrm{d}x + \int_{\partial \Omega} a \frac{\partial u}{\partial n} v \, \mathrm{d}s + \int_{\Omega} f v \, \mathrm{d}s$$

The boundary term can be developed further by noticing that $v \neq 0$ $\partial \Omega_N$,

$$\int_{\partial\Omega} a \frac{\partial u}{\partial n} v \, \mathrm{d}s = \int_{\partial\Omega_N} a \frac{\partial u}{\partial n} v \, \mathrm{d}s,$$

and that on $\partial\Omega_N$, we have the condition $a\frac{\partial u}{\partial n}=-g$, so the term becom

$$-\int_{\partial\Omega_N}gv\,\mathrm{d}s.$$

The variational form is then

$$\int_{\Omega} (\boldsymbol{v} \cdot \nabla u + \alpha u) v \, \mathrm{d}x = -\int_{\Omega} a \nabla u \cdot \nabla v \, \mathrm{d}x - \int_{\partial \Omega_N} g v \, \mathrm{d}s + \int_{\Omega} f v \, \mathrm{d}s$$

Instead of using the integral signs we may use the inner product no

$$(\boldsymbol{v} \cdot \nabla u, v) + (\alpha u, v) = -(a\nabla u, \nabla v) - (g, v)_N + (f, v).$$

The subscript $_N$ in $(g, v)_N$ is a notation for a line or surface integral ov Inserting the u expansion results in

$$\sum_{j \in \mathcal{I}_s} ((\boldsymbol{v} \cdot \nabla \psi_j, \psi_i) + (\alpha \psi_j, \psi_i) + (a \nabla \psi_j, \nabla \psi_i)) c_j =$$

$$(g, \psi_i)_N + (f, \psi_i) - (\boldsymbol{v} \cdot \nabla u_0, \psi_i) + (\alpha u_0, \psi_i) + (a \nabla u_0, \psi_i) +$$

This is a linear system with matrix entries

$$A_{i,j} = (\boldsymbol{v} \cdot \nabla \psi_j, \psi_i) + (\alpha \psi_j, \psi_i) + (a \nabla \psi_j, \nabla \psi_i)$$

and right-hand side entries

$$b_i = (g, \psi_i)_N + (f, \psi_i) - (\boldsymbol{v} \cdot \nabla u_0, \psi_i) + (\alpha u_0, \psi_i) + (a \nabla u_0, \nabla \psi_i)$$

for $i, j \in \mathcal{I}_s$.

In the finite element method, we usually express u_0 in terms of basis fr and restrict i and j to run over the degrees of freedom that are not presc Dirichlet conditions. However, we can also keep all the c_j , $j \in \mathcal{I}_s$, as ur drop the u_0 in the expansion for u, and incorporate all the known c_j v the linear system. This has been explained in detail in the 1D case.

²⁴http://en.wikipedia.org/wiki/Green's_identities

7.1 Transformation to a reference cell in 2D and 3D

le consider an integral of the type

$$\int_{\Omega^{(e)}} a(\mathbf{x}) \nabla \varphi_i \cdot \nabla \varphi_j \, \mathrm{d}x, \tag{201}$$

here the φ_i functions are finite element basis functions in 2D or 3D, defined in 12 physical domain. Suppose we want to calculate this integral over a reference 12 physical domain. Suppose we want to calculate this integral over a reference 13 physical domain. Suppose we want to calculate this integral over a reference 14 physical domain. The mapping between a point \boldsymbol{X} in the reference 15 pordinate system and the corresponding point \boldsymbol{x} in the physical coordinate 15 yet 15 physical domain. The corresponding Jacobian, \boldsymbol{J} , of 15 nis mapping has entries

 $J_{i,j} = \frac{\partial x_j}{\partial X_i} \,.$

The change of variables requires dx to be replaced by $\det J dX$. The derivaves in the ∇ operator in the variational form are with respect to x, which e may denote by ∇_x . The $\varphi_i(x)$ functions in the integral are replaced by cal basis functions $\tilde{\varphi}_r(X)$ so the integral features $\nabla_x \tilde{\varphi}_r(X)$. We readily have $\chi_x \tilde{\varphi}_r(X)$ from formulas for the basis functions in the reference cell, but the esired quantity $\nabla_x \tilde{\varphi}_r(X)$ requires some efforts to compute. All the details are royided below.

Let i = q(e, r) and consider two space dimensions. By the chain rule,

$$\frac{\partial \tilde{\varphi}_r}{\partial X} = \frac{\partial \varphi_i}{\partial X} = \frac{\partial \varphi_i}{\partial x} \frac{\partial x}{\partial X} + \frac{\partial \varphi_i}{\partial y} \frac{\partial y}{\partial X},$$

 $^{\mathrm{nd}}$

$$\frac{\partial \tilde{\varphi}_r}{\partial Y} = \frac{\partial \varphi_i}{\partial Y} = \frac{\partial \varphi_i}{\partial x} \frac{\partial x}{\partial Y} + \frac{\partial \varphi_i}{\partial y} \frac{\partial y}{\partial Y}.$$

/e can write these two equations as a vector equation

$$\begin{bmatrix} \frac{\partial \tilde{\varphi}_r}{\partial X} \\ \frac{\partial \tilde{\varphi}_r}{\partial Y} \end{bmatrix} = \begin{bmatrix} \frac{\partial x}{\partial X} & \frac{\partial y}{\partial X} \\ \frac{\partial x}{\partial Y} & \frac{\partial y}{\partial Y} \end{bmatrix} \begin{bmatrix} \frac{\partial \varphi_i}{\partial x} \\ \frac{\partial \varphi_i}{\partial y} \end{bmatrix}$$

lentifying

$$\nabla_{\boldsymbol{X}} \tilde{\varphi}_r = \begin{bmatrix} \frac{\partial \tilde{\varphi}_r}{\partial X} \\ \frac{\partial \tilde{\varphi}_r}{\partial Y} \end{bmatrix}, \quad J = \begin{bmatrix} \frac{\partial x}{\partial X} & \frac{\partial y}{\partial X} \\ \frac{\partial x}{\partial Y} & \frac{\partial y}{\partial Y} \end{bmatrix}, \quad \nabla_{\boldsymbol{x}} \varphi_r = \begin{bmatrix} \frac{\partial \varphi_i}{\partial x} \\ \frac{\partial \varphi_i}{\partial y} \\ \frac{\partial \varphi_i}{\partial y} \end{bmatrix},$$

e have the relation

$$\nabla_{\mathbf{X}}\tilde{\varphi}_r = J \cdot \nabla_{\mathbf{x}}\varphi_i,$$

hich we can solve with respect to $\nabla_{\boldsymbol{x}}\varphi_i$:

$$\nabla_{\mathbf{x}}\varphi_i = J^{-1} \cdot \nabla_{\mathbf{x}}\tilde{\varphi}_r \,. \tag{202}$$

On the reference cell, $\varphi_i(\boldsymbol{x}) = \tilde{\varphi}_r(\boldsymbol{X})$, so

$$\nabla_{\boldsymbol{x}}\tilde{\varphi}_r(\boldsymbol{X}) = J^{-1}(\boldsymbol{X}) \cdot \nabla_{\boldsymbol{X}}\tilde{\varphi}_r(\boldsymbol{X}).$$

This means that we have the following transformation of the integral physical domain to its counterpart over the reference cell:

$$\int_{\Omega}^{(e)} a(\boldsymbol{x}) \nabla_{\boldsymbol{x}} \varphi_i \cdot \nabla_{\boldsymbol{x}} \varphi_j \, \mathrm{d}\boldsymbol{x} \int_{\tilde{\Omega}^r} a(\boldsymbol{x}(\boldsymbol{X})) (J^{-1} \cdot \nabla_{\boldsymbol{X}} \tilde{\varphi}_r) \cdot (J^{-1} \cdot \nabla \tilde{\varphi}_s) \, \mathrm{det} \, J \, \mathrm{d}\boldsymbol{x}$$

17.2 Numerical integration

Integrals are normally computed by numerical integration rules. Fo dimensional cells, various families of rules exist. All of them are sir what is shown in 1D: $\int f dx \approx \sum_j w_i f(\mathbf{x}_j)$, where w_j are weights and corresponding points.

The file numint.py²⁵ contains the functions quadrature_for_trians and quadrature_for_tetrahedra(n), which returns lists of points and corresponding to integration rules with n points over the reference with vertices (0,0), (1,0), (0,1), and the reference tetrahedron with (0,0,0), (1,0,0), (0,1,0), (0,0,1), respectively. For example, the first to for integration over a triangle have 1 and 3 points:

Rules with 1, 3, 4, and 7 points over the triangle will exactly integrate poly of degree 1, 2, 3, and 4, respectively. In 3D, rules with 1, 4, 5, and 11 over the tetrahedron will exactly integrate polynomials of degree 1, 2, 3 respectively.

17.3 Convenient formulas for P1 elements in 2D

We shall now provide some formulas for piecewise linear φ_i functions a integrals in the physical coordinate system. These formulas make it convecompute with P1 elements without the need to work in the reference coordinate system.

²⁵http://tinyurl.com/nm5587k/fem/numint.py

stem and deal with mappings and Jacobians. A lot of computational and lgorithmic details are hidden by this approach.

Let $\Omega^{(e)}$ be cell number e, and let the three vertices have global vertex umbers I, J, and K. The corresponding coordinates are (x_I, y_I) , (x_J, y_J) , and c_K, y_K . The basis function φ_I over $\Omega^{(e)}$ have the explicit formula

$$\varphi_I(x,y) = \frac{1}{2}\Delta \left(\alpha_I + \beta_I x + \gamma_I y\right), \qquad (205)$$

here

$$\alpha_I = x_J y_K - x_K y_J, \tag{206}$$

$$\beta_I = y_J - y_K, \tag{207}$$

$$\gamma_I = x_K - x_J, \tag{208}$$

$$2\Delta = \det \begin{pmatrix} 1 & x_I & y_I \\ 1 & x_J & y_J \\ 1 & x_K & y_K \end{pmatrix} .$$
 (209)

he quantity Δ is the area of the cell.

The following formula is often convenient when computing element matrices and vectors:

$$\int_{\Omega^{(e)}} \varphi_I^p \varphi_J^q \varphi_K^r dx dy = \frac{p! q! r!}{(p+q+r+2)!} 2\Delta. \tag{210}$$

Note that the q in this formula is not to be mixed with the q(e, r) mapping of egrees of freedom.)

As an example, the element matrix entry $\int_{\Omega^{(e)}} \varphi_I \varphi_J dx$ can be computed y setting p=q=1 and r=0, when $I \neq J$, yielding $\Delta/12$, and p=2 and =r=0, when I=J, resulting in $\Delta/6$. We collect these numbers in a local ement matrix:

$$\frac{\Delta}{12} \left[\begin{array}{ccc} 2 & 1 & 1 \\ 1 & 2 & 1 \\ 1 & 1 & 2 \end{array} \right]$$

The common element matrix entry $\int_{\Omega^{(e)}} \nabla \varphi_I \cdot \nabla \varphi_J \, dx$, arising from a Laplace erm $\nabla^2 u$, can also easily be computed by the formulas above. We have

$$\nabla \varphi_I \cdot \nabla \varphi_J = \frac{\Delta^2}{4} (\beta_I \beta_J + \gamma_I \gamma_J) = \text{const},$$

) that the element matrix entry becomes $\frac{1}{4}\Delta^3(\beta_I\beta_J+\gamma_I\gamma_J)$.

From an implementational point of view, one will work with local vertex umbers r = 0, 1, 2, parameterize the coefficients in the basis functions by r, and ook up vertex coordinates through q(e, r).

Similar formulas exist for integration of P1 elements in 3D.

18 Summary

- When approximating f by $u = \sum_j c_j \varphi_j$, the least squares method Galerkin/projection method give the same result. The interpolatic cation method is simpler and yields different (mostly inferior) res
- Fourier series expansion can be viewed as a least squares or C approximation procedure with sine and cosine functions.
- Basis functions should optimally be orthogonal or almost orthogonal because this gives little round-off errors when solving the linear and the coefficient matrix becomes diagonal or sparse.
- Finite element basis functions are *piecewise* polynomials, norma discontinuous derivatives at the cell boundaries. The basis functions very little, leading to stable numerics and sparse matrices.
- To use the finite element method for differential equations, we Galerkin method or the method of weighted residuals to arrive at tional form. Technically, the differential equation is multiplied b function and integrated over the domain. Second-order derivat integrated by parts to allow for typical finite element basis function have discontinuous derivatives.
- The least squares method is not much used for finite element soldifferential equations of second order, because it then involves second derivatives which cause trouble for basis functions with disconderivatives.
- We have worked with two common finite element terminologies and ated data structures (both are much used, especially the first on the other is more general):
 - 1. elements, nodes, and mapping between local and global node i
 - 2. an extended element concept consisting of cell, vertices, de freedom, local basis functions, geometry mapping, and mapping local and global degrees of freedom
- The meaning of the word "element" is multi-fold: the geometry of element (also known as a cell), the geometry and its basis function information listed under point 2 above.
- One normally computes integrals in the finite element method by element (cell by cell), either in a local reference coordinate sy directly in the physical domain.
- The advantage of working in the reference coordinate system is t mathematical expressions for the basis functions depend on the type only, not the geometry of that element in the physical doma

disadvantage is that a mapping must be used, and derivatives must be transformed from reference to physical coordinates.

- Element contributions to the global linear system are collected in an element matrix and vector, which must be assembled into the global system using the degree of freedom mapping (dof_map) or the node numbering mapping (elements), depending on which terminology that is used.
- Dirichlet conditions, involving prescribed values of u at the boundary, are implemented either via a boundary function that take on the right Dirichlet values, while the basis functions vanish at such boundaries. In the finite element method, one has a general expression for the boundary function, but one can also incorporate Dirichlet conditions in the element matrix and vector or in the global matrix system.
- Neumann conditions, involving prescribed values of the derivative (or flux) of u, are incorporated in boundary terms arising from integrating terms with second-order derivatives by part. Forgetting to account for the boundary terms implies the condition $\partial u/\partial n = 0$ at parts of the boundary where no Dirichlet condition is set.

9 Time-dependent problems

he finite element method is normally used for discretization in space. There re two alternative strategies for performing a discretization in time:

- use finite differences for time derivatives to arrive at a recursive set of spatial problems that can be discretized by the finite element method, or
- discretize in space by finite elements first, and then solve the resulting system of ordinary differential equations (ODEs) by some standard method for ODEs.

le shall exemplify these strategies using a simple diffusion problem

$$\frac{\partial u}{\partial t} = \alpha \nabla^2 u + f(\boldsymbol{x}, t), \qquad \boldsymbol{x} \in \Omega, t \in (0, T], \qquad (211)$$

$$u(\boldsymbol{x},0) = I(\boldsymbol{x}), \qquad \qquad \boldsymbol{x} \in \Omega, \tag{212}$$

$$\frac{\partial u}{\partial n} = 0,$$
 $x \in \partial \Omega, \ t \in (0, T].$ (213)

lere, $u(\boldsymbol{x},t)$ is the unknown function, α is a constant, and $f(\boldsymbol{x},t)$ and I(x) regiven functions. We have assigned the particular boundary condition (213) minimize the details on handling boundary conditions in the finite element nethod.

19.1 Discretization in time by a Forward Euler sche

Time discretization. We can apply a finite difference method in (211). First we need a mesh in time, here taken as uniform with mesh $t_n = n\Delta t, n = 0, 1, \ldots, N_t$. A Forward Euler scheme consists of sa (211) at t_n and approximating the time derivative by a forward di $[D_t^+ u]^n \approx (u^{n+1} - u^n)/\Delta t$. This approximation turns (211) into a diff equation that is discrete in time, but still continuous in space. With difference operator notation we can write the time-discrete problem as

$$[D_t^+ u = \alpha \nabla^2 u + f]^n,$$

for $n=1,2,\ldots,N_t-1$. Writing this equation out in detail and isolat unknown u^{n+1} on the left-hand side, demonstrates that the time-discrete j is a recursive set of problems that are continuous in space:

$$u^{n+1} = u^n + \Delta t \left(\alpha \nabla^2 u^n + f(\boldsymbol{x}, t_n) \right).$$

Given $u^0 = I$, we can use (215) to compute u^1, u^2, \dots, u^{N_t} .

For absolute clarity in the various stages of the discretizations, we in $u_e(\mathbf{x},t)$ as the exact solution of the space-and time-continuous partial differenciation (211) and $u_e^n(\mathbf{x})$ as the time-discrete approximation, arising f finite difference method in time (214). More precisely, u_e fulfills

$$\frac{\partial u_{\mathbf{e}}}{\partial t} = \alpha \nabla^2 u_{\mathbf{e}} + f(\boldsymbol{x}, t),$$

while $u_{\rm e}^{n+1}$, with a superscript, is the solution of the time-discrete equa

$$u_{e}^{n+1} = u_{e}^{n} + \Delta t \left(\alpha \nabla^{2} u_{e}^{n} + f(\boldsymbol{x}, t_{n}) \right).$$

Space discretization. We now introduce a finite element approximation u_{ρ}^{n} and u_{ρ}^{n+1} in (217), where the coefficients depend on the time level:

$$u_{\mathrm{e}}^{n} pprox u^{n} = \sum_{j=0}^{N} c_{j}^{n} \psi_{j}(\boldsymbol{x}),$$
 $u_{\mathrm{e}}^{n+1} pprox u^{n+1} = \sum_{j=0}^{N} c_{j}^{n+1} \psi_{j}(\boldsymbol{x}).$

Note that, as before, N denotes the number of degrees of freedom in the domain. The number of time points is denoted by N_t . We define a spanned by the basis functions $\{\psi_i\}_{i\in\mathcal{I}_t}$.

9.2 Variational forms

weighted residual method with weighting functions w_i can now be formulated. We insert (218) and (219) in (217) to obtain the residual

$$R = u^{n+1} - u^n - \Delta t \left(\alpha \nabla^2 u^n + f(\boldsymbol{x}, t_n) \right).$$

he weighted residual principle,

$$\int_{\Omega} Rw \, \mathrm{d}x = 0, \quad \forall w \in W,$$

sults in

$$\int_{\Omega} \left[u^{n+1} - u^n - \Delta t \left(\alpha \nabla^2 u^n + f(\boldsymbol{x}, t_n) \right) \right] w \, \mathrm{d} x = 0, \quad \forall w \in W.$$

rom now on we use the Galerkin method so W=V. Isolating the unknown $^{n+1}$ on the left-hand side gives

$$\int_{\Omega} u^{n+1} \psi_i \, dx = \int_{\Omega} \left[u^n - \Delta t \left(\alpha \nabla^2 u^n + f(\boldsymbol{x}, t_n) \right) \right] v \, dx, \quad \forall v \in V.$$

As usual in spatial finite element problems involving second-order derivatives, e apply integration by parts on the term $\int (\nabla^2 u^n) v \, dx$:

$$\int_{\Omega} \alpha(\nabla^2 u^n) v \, \mathrm{d}x = -\int_{\Omega} \alpha \nabla u^n \cdot \nabla v \, \mathrm{d}x + \int_{\partial \Omega} \alpha \frac{\partial u^n}{\partial n} v \, \mathrm{d}x.$$

he last term vanishes because we have the Neumann condition $\partial u^n/\partial n = 0$ for $\ln n$. Our discrete problem in space and time then reads

$$\int_{\Omega} u^{n+1} v \, dx = \int_{\Omega} u^n v dx - \Delta t \int_{\Omega} \alpha \nabla u^n \cdot \nabla v \, dx + \Delta t \int_{\Omega} f^n v \, dx, \quad \forall \boldsymbol{v} \in V.$$
(220)

his is the variational formulation of our recursive set of spatial problems.

Nonzero Dirichlet boundary conditions.

As in stationary problems, we can introduce a boundary function B(x,t) to take care of nonzero Dirichlet conditions:

$$u_{\rm e}^n \approx u^n = B(\boldsymbol{x}, t_n) + \sum_{j=0}^N c_j^n \psi_j(\boldsymbol{x}),$$

 $u_{\rm e}^{n+1} \approx u^{n+1} = B(\boldsymbol{x}, t_{n+1}) + \sum_{j=0}^N c_j^{n+1} \psi_j(\boldsymbol{x}).$

19.3 Simplified notation for the solution at recent levels

In a program it is only necessary to store u^{n+1} and u^n at the same tire therefore drop the n index in programs and work with two functions: u f the new unknown, and u_1 for u^n , the solution at the previous time levis also convenient in the mathematics to maximize the correspondent he code. From now on u_1 means the discrete unknown at the previous level (u^n) and u represents the discrete unknown at the new time level Equation (220) with this new naming convention is expressed as

$$\int_{\Omega} uv dx = \int_{\Omega} u_1 v dx - \Delta t \int_{\Omega} \alpha \nabla u_1 \cdot \nabla v \, dx + \Delta t \int_{\Omega} f^n v \, dx.$$

This variational form can alternatively be expressed by the inner prodution:

$$(u,v) = (u_1,v) - \Delta t(\alpha \nabla u_1, \nabla v) + (f^n,v).$$

19.4 Deriving the linear systems

To derive the equations for the new unknown coefficients c_j^{n+1} , now jus c_j , we insert

$$u = \sum_{j=0}^{N} c_j \psi_j(x), \quad u_1 = \sum_{j=0}^{N} c_{1,j} \psi_j(x)$$

in (223) or (224), let the equation hold for all $v=\psi,\,i=0,\ldots,\!{\rm N},$ and o terms as matrix-vector products:

$$\sum_{j=0}^{N} (\psi_i, \psi_j) c_j = \sum_{j=0}^{N} (\psi_i, \psi_j) c_{1,j} - \Delta t \sum_{j=0}^{N} (\nabla \psi_i, \alpha \nabla \psi_j) c_{1,j} + (f^n, \psi_i), \quad i = 0$$

This is a linear system $\sum_{i} A_{i,j} c_j = b_i$ with

$$A_{i,j} = (\psi_i, \psi_j)$$

nd

$$b_{i} = \sum_{j=0}^{N} (\psi_{i}, \psi_{j}) c_{1,j} - \Delta t \sum_{j=0}^{N} (\nabla \psi_{i}, \alpha \nabla \psi_{j}) c_{1,j} + (f^{n}, \psi_{i}).$$

It is instructive and convenient for implementations to write the linear system **n** the form

$$Mc = Mc_1 - \Delta t K c_1 + f, \tag{226}$$

here

$$\begin{split} M &= \{M_{i,j}\}, \quad M_{i,j} = (\psi_i, \psi_j), \quad i, j \in \mathcal{I}_s, \\ K &= \{K_{i,j}\}, \quad K_{i,j} = (\nabla \psi_i, \alpha \nabla \psi_j), \quad i, j \in \mathcal{I}_s, \\ f &= \{(f(\boldsymbol{x}, t_n), \psi_i)\}_{i \in \mathcal{I}_s}, \\ c &= \{c_i\}_{i \in \mathcal{I}_s}, \\ c_1 &= \{c_{1,i}\}_{i \in \mathcal{I}_s}. \end{split}$$

We realize that M is the matrix arising from a term with the zero-th derivative f u, and called the mass matrix, while K is the matrix arising from a Laplace $\operatorname{rm} \nabla^2 u$. The K matrix is often known as the $\operatorname{stiffness}$ matrix. (The terms ass and stiffness stem from the early days of finite elements when applications $\mathfrak m$ vibrating structures dominated. The mass matrix arises from the mass times coeleration term in Newton's second law, while the stiffness matrix arises from the elastic forces in that law. The mass and stiffness matrix appearing in a iffusion have slightly different mathematical formulas.)

temark. The mathematical symbol f has two meanings, either the function (x,t) in the PDE or the f vector in the linear system to be solved at each time vel. The symbol u also has different meanings, basically the unknown in the DE or the finite element function representing the unknown at a time level. he actual meaning should be evident from the context.

9.5 Computational algorithm

/e observe that M and K can be precomputed so that we can avoid computing ne matrix entries at every time level. Instead, some matrix-vector multiplications ill produce the linear system to be solved. The computational algorithm has ne following steps:

- 1. Compute M and K.
- 2. Initialize u^0 by interpolation or projection

- 3. For $n = 1, 2, ..., N_t$:
 - (a) compute $b = Mc_1 \Delta t K c_1 + f$
 - (b) solve Mc = b
 - (c) set $c_1 = c$

In case of finite element basis functions, interpolation of the initial coat the nodes means $c_{1,j} = I(\boldsymbol{x}_j)$. Otherwise one has to solve the linear $\sum_j \psi_j(x_i)c_j = I(x_i)$, where \boldsymbol{x}_j denotes an interpolation point. Projec Galerkin's method) implies solving a linear system with M as coefficient: $\sum_j M_{i,j}c_{1,j} = (I,\psi_i), i \in \mathcal{I}_s$.

19.6 Comparing P1 elements with the finite different method

We can compute the M and K matrices using P1 elements in 1D. A mesh on [0, L] is introduced for this purpose. Since the boundary condit solely of Neumann type in this sample problem, we have no restrictions basis functions ψ_i and can simply choose $\psi_i = \varphi_i$, $i = 0, ..., N = N_n$.

From Section 13.2 or 13.4 we have that the K matrix is the same as from the finite difference method: $h[D_xD_xu]_i^n$, while from Section 5.2 v that M can be interpreted as the finite difference approximation $[u+\frac{1}{6}h^2L$ (times h). The equation system Mc=b in the algorithm is therefore eq to the finite difference scheme

$$[D_t^+(u + \frac{1}{6}h^2D_xD_xu) = \alpha D_xD_xu + f]_i^n.$$

(More precisely, Mc = b divided by h gives the equation above.)

Lumping the mass matrix. By applying Trapezoidal integration turn M into a diagonal matrix with $(h/2, h, \ldots, h, h/2)$ on the diagonal there is no need to solve a linear system at each time level, and the finite scheme becomes identical to a standard finite difference method

$$[D_t^+ u = \alpha D_x D_x u + f]_i^n.$$

The Trapezoidal integration is not as accurate as exact integrating introduces therefore an error. Whether this error has a good or bad in on the overall numerical method is not immediately obvious, and is an adetail in Section 19.10. The effect of the error is at least not more seven what is produced by the finite difference method.

Making M diagonal is usually referred to as *lumping the mass matrix* is an alternative method to using an integration rule based on the node one can sum the entries in each row, place the sum on the diagonal, and other entries in the row equal to zero. For P1 elements the methods of late mass matrix give the same result.

9.7 Discretization in time by a Backward Euler scheme

'ime discretization. The Backward Euler scheme in time applied to our iffusion problem can be expressed as follows using the finite difference operator otation:

$$[D_t^- u = \alpha \nabla^2 u + f(\boldsymbol{x}, t)]^n.$$

/ritten out, and collecting the unknown u^n on the left-hand side and all the nown terms on the right-hand side, the time-discrete differential equation ecomes

$$u_{\mathrm{e}}^{n} - \Delta t \left(\alpha \nabla^{2} u_{\mathrm{e}}^{n} + f(\boldsymbol{x}, t_{n}) \right) = u_{\mathrm{e}}^{n-1}. \tag{229}$$

quation (229) can compute $u_{\rm e}^1, u_{\rm e}^2, \ldots, u_{\rm e}^{N_t}$, if we have a start $u_{\rm e}^0 = I$ from the itial condition. However, (229) is a partial differential equation in space and eeds a solution method based on discretization in space. For this purpose we se an expansion as in (218)-(219).

'ariational forms. Inserting (218)-(219) in (229), multiplying by ψ_i (or $\in V$), and integrating by parts, as we did in the Forward Euler case, results in 12 variational form

$$\int_{\Omega} (u^n v + \Delta t \alpha \nabla u^n \cdot \nabla v) \, dx = \int_{\Omega} u^{n-1} v \, dx - \Delta t \int_{\Omega} f^n v \, dx, \quad \forall v \in V. \quad (230)$$

xpressed with u as u^n and u_1 as u^{n-1} , this becomes

$$\int_{\Omega} (uv + \Delta t \alpha \nabla u \cdot \nabla v) \, dx = \int_{\Omega} u_1 v \, dx + \Delta t \int_{\Omega} f^n v \, dx, \qquad (231)$$

r with the more compact inner product notation,

$$(u,v) + \Delta t(\alpha \nabla u, \nabla v) = (u_1, v) + \Delta t(f^n, v).$$
(232)

inear systems. Inserting $u = \sum_j c_j \psi_i$ and $u_1 = \sum_j c_{1,j} \psi_i$, and choosing v be the basis functions $\psi_i \in V$, i = 0, ..., N, together with doing some algebra, and to the following linear system to be solved at each time level:

$$(M + \Delta t K)c = Mc_1 + f, \tag{233}$$

here M, K, and f are as in the Forward Euler case. This time we really have a solve a linear system at each time level. The computational algorithm goes as ollows.

- 1. Compute M, K, and $A = M + \Delta t K$
- 2. Initialize u^0 by interpolation or projection

- 3. For $n = 1, 2, \dots, N_t$:
 - (a) compute $b = Mc_1 + f$
 - (b) solve Ac = b
 - (c) set $c_1 = c$

In case of finite element basis functions, interpolation of the initial coat the nodes means $c_{1,j} = I(\boldsymbol{x}_j)$. Otherwise one has to solve the linear $\sum_j \psi_j(x_i)c_j = I(x_i)$, where \boldsymbol{x}_j denotes an interpolation point. Projec Galerkin's method) implies solving a linear system with M as coefficient : $\sum_j M_{i,j}c_{1,j} = (I,\psi_i)$, $i \in \mathcal{I}_s$.

We know what kind of finite difference operators the M and K r correspond to (after dividing by h), so (233) can be interpreted as the finite difference method:

$$[D_t^-(u + \frac{1}{6}h^2D_xD_xu) = \alpha D_xD_xu + f]_i^n.$$

The mass matrix M can be lumped, as explained in Section 19.6, a the linear system arising from the finite element method with P1 e corresponds to a plain Backward Euler finite difference method for the equation:

$$[D_t^- u = \alpha D_x D_x u + f]_i^n.$$

19.8 Dirichlet boundary conditions

Suppose now that the boundary condition (213) is replaced by a mixed N and Dirichlet condition,

$$u(\boldsymbol{x},t) = u_0(\boldsymbol{x},t), \qquad \qquad \boldsymbol{x} \in \partial \Omega_D, \ -\alpha \frac{\partial}{\partial n} u(\boldsymbol{x},t) = g(\boldsymbol{x},t), \qquad \qquad \boldsymbol{x} \in \partial \Omega_N.$$

Using a Forward Euler discretization in time, the variational form at level becomes

$$\int_{\Omega} u^{n+1} v \, \mathrm{d}x = \int_{\Omega} (u^n - \Delta t \alpha \nabla u^n \cdot \nabla v) \, \mathrm{d}x - \Delta t \int_{\partial \Omega_N} g v \, \mathrm{d}s, \quad \forall v \in V$$

Boundary function. The Dirichlet condition $u = u_0$ at $\partial \Omega_D$ can be rated through a boundary function $B(\mathbf{x}) = u_0(\mathbf{x})$ and demanding that $\partial \Omega_D$. The expansion for u^n is written as

$$u^n(\mathbf{x}) = u_0(\mathbf{x}, t_n) + \sum_{j \in \mathcal{I}_s} c_j^n \psi_j(\mathbf{x}).$$

iserting this expansion in the variational formulation and letting it hold for all asis functions ψ_i leads to the linear system

$$\sum_{i \in \mathcal{I}_s} \left(\int_{\Omega} \psi_i \psi_j \, \mathrm{d}x \right) c_j^{n+1} = \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \left(\psi_i \psi_j - \Delta t \alpha \nabla \psi_i \cdot \nabla \psi_j \right) \, \mathrm{d}x \right) c_j^n - \int_{\Omega} \left(u_0(\boldsymbol{x}, t_{n+1}) - u_0(\boldsymbol{x}, t_n) + \Delta t \alpha \nabla u_0(\boldsymbol{x}, t_n) \cdot \nabla \psi_i \right) \, \mathrm{d}x + \Delta t \int_{\Omega} f \psi_i \, \mathrm{d}x - \Delta t \int_{\partial \Omega_N} g \psi_i \, \mathrm{d}s, \quad i \in \mathcal{I}_s.$$

1 the following, we adopt the convention that the unknowns c_j^{n+1} are written c_j , while the known c_j^n from the previous time level are denoted by $c_{1,j}$.

inite element basis functions. When using finite elements, each basis inction φ_i is associated with a node x_i . We have a collection of nodes $\{x_i\}_{i\in I_b}$ in the boundary $\partial\Omega_D$. Suppose U_k^n is the known Dirichlet value at x_k at time Ω_k^n ($U_k^n = u_0(x_k, t_n)$). The appropriate boundary function is then

$$B(\boldsymbol{x},t_n) = \sum_{j \in I_h} U_j^n \varphi_j.$$

he unknown coefficients c_j are associated with the rest of the nodes, which are numbers $\nu(i)$, $i \in \mathcal{I}_s = \{0, \dots, N\}$. The basis functions for V are chosen as $i = \varphi_{\nu(i)}$, $i \in \mathcal{I}_s$, and all of these vanish at the boundary nodes as they should he expansion for u^{n+1} and u^n become

$$u^{n} = \sum_{j \in I_{b}} U_{j}^{n} \varphi_{j} + \sum_{j \in \mathcal{I}_{s}} c_{1,j} \varphi_{\nu(j)},$$

$$u^{n+1} = \sum_{j \in I_{b}} U_{j}^{n+1} \varphi_{j} + \sum_{j \in \mathcal{I}_{s}} c_{j} \varphi_{\nu(j)}.$$

he equations for the unknown coefficients c_i become

$$\sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \varphi_i \varphi_j \, \mathrm{d}x \right) c_j = \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \left(\varphi_i \varphi_j - \Delta t \alpha \nabla \varphi_i \cdot \nabla \varphi_j \right) \, \mathrm{d}x \right) c_{1,j} - \sum_{j \in I_b} \int_{\Omega} \left(\varphi_i \varphi_j (U_j^{n+1} - U_j^n) + \Delta t \alpha \nabla \varphi_i \cdot \nabla \varphi_j U_j^n \right) \, \mathrm{d}x + \Delta t \int_{\Omega} f \varphi_i \, \mathrm{d}x - \Delta t \int_{\partial \Omega_N} g \varphi_i \, \mathrm{d}s, \quad i \in \mathcal{I}_s.$$

Modification of the linear system. Instead of introducing a befunction B we can work with basis functions associated with all the no incorporate the Dirichlet conditions by modifying the linear system. Let the index set that counts all the nodes: $\{0, 1, \ldots, N = N_n\}$. The expan u^n is then $\sum_{j \in \mathcal{I}_s} c_j^n \varphi_j$ and the variational form becomes

$$\sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \varphi_i \varphi_j \, \mathrm{d}x \right) c_j = \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \left(\varphi_i \varphi_j - \Delta t \alpha \nabla \varphi_i \cdot \nabla \varphi_j \right) \, \mathrm{d}x \right) c_1,$$
$$- \Delta t \int_{\Omega} f \varphi_i \, \mathrm{d}x - \Delta t \int_{\partial \Omega_N} g \varphi_i \, \mathrm{d}s.$$

We introduce the matrices M and K with entries $M_{i,j} = \int_{\Omega} \varphi_i \varphi_j \, dx$ and $\int_{\Omega} \alpha \nabla \varphi_i \cdot \nabla \varphi_j \, dx$, respectively. In addition, we define the vectors c, c_1 with entries $c_i, c_{1,i}$, and $\int_{\Omega} f \varphi_i \, dx - \int_{\partial \Omega_N} g \varphi_i \, ds$. The equation system c_i be written as

$$Mc = Mc_1 - \Delta t K c_1 + \Delta t f.$$

When M, K, and b are assembled without paying attention to Dirichlet be conditions, we need to replace equation k by $c_k = U_k$ for k corresponds to all boundary nodes $(k \in I_b)$. The modification of M consists in $M_{k,j} = 0$, $j \in \mathcal{I}_s$, and the $M_{k,k} = 1$. Alternatively, a modification preserves the symmetry of M can be applied. At each time level on $b = Mc_1 - \Delta t K c_1 + \Delta t f$ and sets $b_k = U_k^{n+1}$, $k \in I_b$, and solves the Mc = b.

In case of a Backward Euler method, the system becomes (233). write the system as Ac = b, with $A = M + \Delta t K$ and $b = Mc_1 + f$. I and K needs to be modified because of Dirichlet boundary conditions, diagonal entries in K should be set to zero and those in M to unity. way, $A_{k,k} = 1$. The right-hand side must read $b_k = U_k^n$ for $k \in I_b$ (assun unknown is sought at time level t_n).

19.9 Example: Oscillating Dirichlet boundary cond

We shall address the one-dimensional initial-boundary value problem

$$u_t = (\alpha u_x)_x + f,$$
 $x \in \Omega = [0, L], t \in (0, T],$
 $u(x, 0) = 0,$ $x \in \Omega,$
 $u(0, t) = a \sin \omega t,$ $t \in (0, T],$
 $u_x(L, t) = 0,$ $t \in (0, T].$

A physical interpretation may be that u is the temperature deviation a constant mean temperature in a body Ω that is subject to an osci temperature (e.g., day and night, or seasonal, variations) at x = 0.

We use a Backward Euler scheme in time and P1 elements of constant length in space. Incorporation of the Dirichlet condition at x=0 through modifying ne linear system at each time level means that we carry out the computations s explained in Section 19.7 and get a system (233). The M and K matrices omputed without paying attention to Dirichlet boundary conditions become

he right-hand side of the variational form contains Mc_1 since there is no source $\operatorname{rm}(f)$ and no boundary term from the integration by parts $(u_x=0 \text{ at } x=L \text{ nd})$ we compute as if $u_x=0 \text{ at } x=0 \text{ too}$. We must incorporate the Dirichlet oundary condition $c_0=a\sin\omega t_n$ by ensuring that this is the first equation in the linear system. To this end, the first row in K and M are set to zero, but the diagonal entry $M_{0,0}$ is set to 1. The right-hand side is $b=Mc_1$, and we set $c_0=a\sin\omega t_n$. Note that in this approach, $c_0=a\sin\omega t_n$, and $c_0=a\sin\omega t_n$ are each node in the mesh. We can write the complete linear system as

$$c_0 = a \sin \omega t_n,$$

$$\frac{h}{6}(c_{i-1} + 4c_i + c_{i+1}) + \Delta t \frac{\alpha}{h}(-c_{i-1} + 2c_i + c_{i+1}) = \frac{h}{6}(c_{1,i-1} + 4c_{1,i} + c_1)$$

$$i = 1, \dots, N_n - 1,$$

$$\frac{h}{6}(c_{i-1} + 2c_i) + \Delta t \frac{\alpha}{h}(-c_{i-1} + c_i) = \frac{h}{6}(c_{1,i-1} + 2c_{1,i}),$$

The Dirichlet boundary condition can alternatively be implemented a boundary function $B(x,t) = a \sin \omega t \varphi_0(x)$:

$$u^{n}(x) = a \sin \omega t_{n} \varphi_{0}(x) + \sum_{j \in \mathcal{I}_{s}} c_{j} \varphi_{\nu(j)}(x), \quad \nu(j) = j + 1.$$

Now, $N = N_n - 1$ and the c vector contains values of u at nodes 1, 2, The right-hand side gets a contribution

$$\int_0^L \left(a(\sin\omega t_n - \sin\omega t_{n-1})\varphi_0\varphi_i - \Delta t\alpha a\sin\omega t_n\nabla\varphi_0\cdot\nabla\varphi_i\right)\,\mathrm{d}x\,.$$

19.10 Analysis of the discrete equations

The diffusion equation $u_t = \alpha u_{xx}$ allows a (Fourier) wave compone $\exp(\beta t + ikx)$ as solution if $\beta = -\alpha k^2$, which follows from inserting the component in the equation. The exact wave component can alternative written as

$$u = A_e^n e^{ikx}, \quad A_e = e^{-\alpha k^2 \Delta t}.$$

Many numerical schemes for the diffusion equation has a similar wave cor as solution:

$$u_q^n = A^n e^{ikx},$$

where is an amplification factor to be calculated by inserting (252) in the We introduce x = qh, or $x = q\Delta x$ to align the notation with that fre used in finite difference methods.

A convenient start of the calculations is to establish some results for finite difference operators acting on

$$u_q^n = A^n e^{ikq\Delta x}$$

$$\begin{split} [D_t^+ A^n e^{ikq\Delta x}]^n &= A^n e^{ikq\Delta x} \frac{A-1}{\Delta t}, \\ [D_t^- A^n e^{ikq\Delta x}]^n &= A^n e^{ikq\Delta x} \frac{1-A^{-1}}{\Delta t}, \\ [D_t A^n e^{ikq\Delta x}]^{n+\frac{1}{2}} &= A^{n+\frac{1}{2}} e^{ikq\Delta x} \frac{A^{\frac{1}{2}} - A^{-\frac{1}{2}}}{\Delta t} = A^n e^{ikq\Delta x} \frac{A-1}{\Delta t}, \\ [D_x D_x A^n e^{ikq\Delta x}]_q &= -A^n \frac{4}{\Delta x^2} \sin^2 \left(\frac{k\Delta x}{2}\right). \end{split}$$

orward Euler discretization. We insert (252) in the Forward Euler scheme ith P1 elements in space and f = 0 (this type of analysis can only be carried at if f = 0),

$$[D_t^+(u + \frac{1}{6}h^2D_xD_xu) = \alpha D_xD_xu]_q^n.$$
 (253)

le have

$$D_t^+ D_x D_x A e^{ikx}]_q^n = [D_t^+ A]^n [D_x D_x e^{ikx}]_q = -A^n e^{ikp\Delta x} \frac{A-1}{\Delta t} \frac{4}{\Delta x^2} \sin^2(\frac{k\Delta x}{2}).$$

he term $[D_t^+Ae^{ikx} + \frac{1}{6}\Delta x^2D_t^+D_xD_xAe^{ikx}]_a^n$ then reduces to

$$\frac{A-1}{\Delta t} - \frac{1}{6}\Delta x^2 \frac{A-1}{\Delta t} \frac{4}{\Delta x^2} \sin^2(\frac{k\Delta x}{2}),$$

 $\frac{A-1}{\Delta t} \left(1 - \frac{2}{3} \sin^2(k\Delta x/2) \right) .$

itroducing $p = k\Delta x/2$ and $C = \alpha \Delta t/\Delta x^2$, the complete scheme becomes

$$(A-1)\left(1 - \frac{2}{3}\sin^2 p\right) = -4C\sin^2 p,$$

om which we find A to be

$$A = 1 - 4C \frac{\sin^2 p}{1 - \frac{2}{3}\sin^2 p} \,.$$

How does this A change the stability criterion compared to the Forward uler finite difference scheme and centered differences in space? The stability iterion is $|A| \leq 1$, which here implies $A \leq 1$ and $A \geq -1$. The former is always ilfilled, while the latter leads to

$$4C\frac{\sin^2 p}{1+\frac{2}{3}\sin^2 p} \le 2.$$

The factor $\sin^2 p/(1-\frac{2}{3}\sin^2 p)$ can be plotted for $p \in [0,\pi/2]$, and the m value goes to 3 as $p \to \pi/2$. The worst case for stability therefore occurs shortest possible wave, $p = \pi/2$, and the stability criterion becomes

$$C \le \frac{1}{6} \quad \Rightarrow \quad \Delta t \le \frac{\Delta x^2}{6\alpha},$$

which is a factor 1/3 worse than for the standard Forward Euler finite dimethod for the diffusion equation, which demands $C \leq 1/2$. Lumping t matrix will, however, recover the finite difference method and therefor $C \leq 1/2$ for stability.

Backward Euler discretization. We can use the same approach an (252) in the Backward Euler scheme with P1 elements in space and f =

$$[D_t^-(u + \frac{1}{6}h^2D_xD_xu) = \alpha D_xD_xu]_i^n.$$

Similar calculations as in the Forward Euler case lead to

$$(1 - A^{-1})\left(1 - \frac{2}{3}\sin^2 p\right) = -4C\sin^2 p,$$

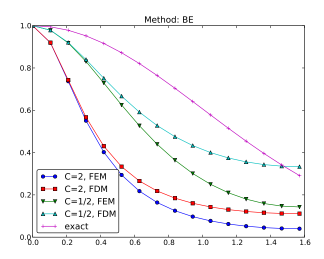
and hence

$$A = \left(1 + 4C \frac{\sin^2 p}{1 - \frac{2}{3}\sin^2 p}\right)^{-1}.$$

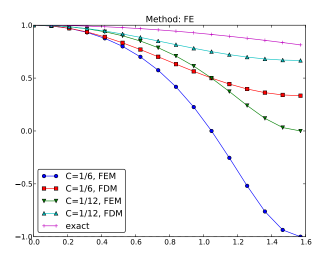
Comparing amplification factors. It is of interest to compare A are functions of p for some C values. Figure 48 display the amplification for the Backward Euler scheme corresponding a coarse mesh with C= mesh at the stability limit of the Forward Euler scheme in the finite di method, C=1/2. Figures 49 and 50 shows how the accuracy increas lower C values for both the Forward Euler and Backward schemes, resp. The striking fact, however, is that the accuracy of the finite element is significantly less than the finite difference method for the same valumping the mass matrix to recover the numerical amplification factor finite difference method is therefore a good idea in this problem.

Remaining tasks:

- Taylor expansion of the error in the amplification factor $A_{\rm e}-A$
- Taylor expansion of the error $e = (A_e^n A^n)e^{ikx}$
- L^2 norm of e



igure 48: Comparison of coarse-mesh amplification factors for Backward Euler iscretization of a 1D diffusion equation.



igure 49: Comparison of fine-mesh amplification factors for Forward Euler iscretization of a 1D diffusion equation.

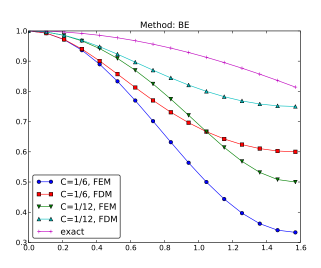


Figure 50: Comparison of fine-mesh amplification factors for Backwar discretization of a 1D diffusion equation.

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30 Systems of differential equations

Inny mathematical models involve m+1 unknown functions governed by a ystem of m+1 differential equations. In abstract form we may denote the nknowns by $u^{(0)}, \ldots, u^{(m)}$ and write the governing equations as

$$\mathcal{L}_0(u^{(0)}, \dots, u^{(m)}) = 0,$$

 \vdots
 $\mathcal{L}_m(u^{(0)}, \dots, u^{(m)}) = 0,$

here \mathcal{L}_i is some differential operator defining differential equation number i.

0.1 Variational forms

here are basically two ways of formulating a variational form for a system of ifferential equations. The first method treats each equation independently as scalar equation, while the other method views the total system as a vector quation with a vector function as unknown.

Let us start with the one equation at a time approach. We multiply equation umber i by some test function $v^{(i)} \in V^{(i)}$ and integrate over the domain:

$$\int_{\Omega} \mathcal{L}^{(0)}(u^{(0)}, \dots, u^{(m)}) v^{(0)} \, \mathrm{d}x = 0, \tag{256}$$

$$\int_{\Omega} \mathcal{L}^{(m)}(u^{(0)}, \dots, u^{(m)}) v^{(m)} \, \mathrm{d}x = 0.$$
 (258)

erms with second-order derivatives may be integrated by parts, with Neumann onditions inserted in boundary integrals. Let

$$V^{(i)} = \operatorname{span}\{\psi_0^{(i)}, \dots, \psi_{N_i}^{(i)}\},\$$

ich that

$$u^{(i)} = B^{(i)}(\boldsymbol{x}) + \sum_{j=0}^{N_i} c_j^{(i)} \psi_j^{(i)}(\boldsymbol{x}),$$

here $B^{(i)}$ is a boundary function to handle nonzero Dirichlet conditions. Observe at different unknowns live in different spaces with different basis functions and umbers of degrees of freedom.

From the m equations in the variational forms we can derive m coupled ystems of algebraic equations for the $\prod_{i=0}^{m} N_i$ unknown coefficients $c_j^{(i)}$, $j = \dots, N_i, i = 0, \dots, m$.

The alternative method for deriving a variational form for a sy differential equations introduces a vector of unknown functions

$$\boldsymbol{u} = (u^{(0)}, \dots, u^{(m)}),$$

a vector of test functions

$$\mathbf{v} = (u^{(0)}, \dots, u^{(m)})$$

with

$$\boldsymbol{u}, \boldsymbol{v} \in \boldsymbol{V} = V^{(0)} \times \cdots \times V^{(m)}$$
.

With nonzero Dirichlet conditions, we have a vector $\mathbf{B} = (B^{(0)}, \dots, B^{(n)})$ boundary functions and then it is $\mathbf{u} - \mathbf{B}$ that lies in \mathbf{V} , not \mathbf{u} itself.

The governing system of differential equations is written

$$\mathcal{L}(\mathbf{u}) = 0$$
,

where

$$\mathcal{L}(oldsymbol{u}) = (\mathcal{L}^{(0)}(oldsymbol{u}), \ldots, \mathcal{L}^{(m)}(oldsymbol{u}))$$

The variational form is derived by taking the inner product of the vequations and the test function vector:

$$\int_{\Omega} \mathcal{L}(\boldsymbol{u}) \cdot \boldsymbol{v} = 0 \quad \forall \boldsymbol{v} \in \boldsymbol{V}.$$

Observe that (259) is one scalar equation. To derive systems of a equations for the unknown coefficients in the expansions of the unknown tions, one chooses m linearly independent \boldsymbol{v} vectors to generate m independent forms (259). The particular choice $\boldsymbol{v}=(v^{(0)},0,\ldots,0)$: (256), $\boldsymbol{v}=(0,\ldots,0,v^{(m)})$ recovers (258), and $\boldsymbol{v}=(0,\ldots,0,v^{(i)},0,\ldots,0)$ the variational form number i, $\int_{\Omega} \mathcal{L}^{(i)} v^{(i)} \, \mathrm{d} x = 0$, in (256)-(258).

20.2 A worked example

We now consider a specific system of two partial differential equations space dimensions:

$$\mu \nabla^2 w = -\beta,$$

$$\kappa \nabla^2 T = -\mu ||\nabla w||^2.$$

The unknown functions w(x,y) and T(x,y) are defined in a domain (μ, β, μ) and (κ) are given constants. The norm in (261) is the standard Eunorm:

$$||\nabla w||^2 = \nabla w \cdot \nabla w = w_x^2 + w_y^2.$$

The boundary conditions associated with (260)-(261) are w = 0 on $\partial\Omega$ and $' = T_0$ on $\partial\Omega$. Each of the equations (260) and (261) need one condition at ach point on the boundary.

The system (260)-(261) arises from fluid flow in a straight pipe, with the z axis the direction of the pipe. The domain Ω is a cross section of the pipe, w is the elocity in the z direction, μ is the viscosity of the fluid, β is the pressure gradient long the pipe, T is the temperature, and κ is the heat conduction coefficient f the fluid. The equation (260) comes from the Navier-Stokes equations, and (261) follows from the energy equation. The term $-\mu ||\nabla w||^2$ models heating of the fluid due to internal friction.

Observe that the system (260)-(261) has only a one-way coupling: T depends n w, but w does not depend on T, because we can solve (260) with respect w and then (261) with respect to w. Some may argue that this is not a real system of PDEs, but just two scalar PDEs. Nevertheless, the one-way oupling is convenient when comparing different variational forms and different nplementations.

0.3 Identical function spaces for the unknowns

et us first apply the same function space V for w and T (or more precisely, $v \in V$ and $T - T_0 \in V$). With

$$V = \operatorname{span}\{\psi_0(x, y), \dots, \psi_N(x, y)\},\$$

e write

$$w = \sum_{j=0}^{N} c_j^{(w)} \psi_j, \quad T = T_0 + \sum_{j=0}^{N} c_j^{(T)} \psi_j.$$
 (262)

ote that w and T in (260)-(261) denote the exact solution of the PDEs, while v and v (262) are the discrete functions that approximate the exact solution. Should be clear from the context whether a symbol means the exact or pproximate solution, but when we need both at the same time, we use a abscript v to denote the exact solution.

'ariational form of each individual PDE. Inserting the expansions (262) 1 the governing PDEs, results in a residual in each equation,

$$R_w = \mu \nabla^2 w + \beta, \tag{263}$$

$$R_T = \kappa \nabla^2 T + \mu ||\nabla w||^2. \tag{264}$$

Galerkin method demands R_w and R_T do be orthogonal to V:

$$\int_{\Omega} R_w v \, \mathrm{d}x = 0 \quad \forall v \in V,$$

$$\int_{\Omega} R_T v \, \mathrm{d}x = 0 \quad \forall v \in V.$$

Because of the Dirichlet conditions, v=0 on $\partial\Omega$. We integrate the terms by parts and note that the boundary terms vanish since v=0 or

$$\int_{\Omega} \mu \nabla w \cdot \nabla v \, dx = \int_{\Omega} \beta v \, dx \quad \forall v \in V,$$
$$\int_{\Omega} \kappa \nabla T \cdot \nabla v \, dx = \int_{\Omega} \mu \nabla w \cdot \nabla w \, v \, dx \quad \forall v \in V.$$

Compound scalar variational form. The alternative way of derivariational from is to introduce a test vector function $\mathbf{v} \in \mathbf{V} = V \times V$ at the inner product of \mathbf{v} and the residuals, integrated over the domain:

$$\int_{\Omega} (R_w, R_T) \cdot \boldsymbol{v} \, \mathrm{d}x = 0 \quad \forall \boldsymbol{v} \in \boldsymbol{V}.$$

With $\mathbf{v} = (v_0, v_1)$ we get

$$\int_{\Omega} (R_w v_0 + R_T v_1) \, \mathrm{d}x = 0 \quad \forall \boldsymbol{v} \in \boldsymbol{V} \,.$$

Integrating the Laplace terms by parts results in

$$\int_{\Omega} (\mu \nabla w \cdot \nabla v_0 + \kappa \nabla T \cdot \nabla v_1) \, dx = \int_{\Omega} (\beta v_0 + \mu \nabla w \cdot \nabla w \, v_1) \, dx, \quad \forall \boldsymbol{v} \in \boldsymbol{V}$$

Choosing $v_0 = v$ and $v_1 = 0$ gives the variational form (265), while $v_0 : v_1 = v$ gives (266).

With the inner product notation, $(p,q) = \int_{\Omega} pq \, dx$, we can alternative (265) and (266) as

$$(\mu \nabla w, \nabla v) = (\beta, v) \quad \forall v \in V,$$

$$(\kappa \nabla T, \nabla v) = (\mu \nabla w \cdot \nabla w, v) \quad \forall v \in V.$$

or since μ and κ are considered constant,

$$\mu(\nabla w, \nabla v) = (\beta, v) \quad \forall v \in V,$$

$$\kappa(\nabla T, \nabla v) = \mu(\nabla w \cdot \nabla w, v) \quad \forall v \in V.$$

Decoupled linear systems. The linear systems governing the coefficients $c_j^{(w)}$ and $c_j^{(T)}$, $j=0,\ldots,N$, are derived by inserting the expansions (262) in (265) and (266), and choosing $v=\psi_i$ for $i=0,\ldots,N$. The result becomes

$$\sum_{j=0}^{N} A_{i,j}^{(w)} c_j^{(w)} = b_i^{(w)}, \quad i = 0, \dots, N,$$
(270)

$$\sum_{i=0}^{N} A_{i,j}^{(T)} c_j^{(T)} = b_i^{(T)}, \quad i = 0, \dots, N,$$
(271)

$$A_{i,j}^{(w)} = \mu(\nabla \psi_j, \nabla \psi_i), \tag{272}$$

$$b_i^{(w)} = (\beta, \psi_i), \tag{273}$$

$$A_{i,j}^{(T)} = \kappa(\nabla \psi_j, \nabla \psi_i), \tag{274}$$

$$b_i^{(T)} = \mu((\sum_j c_j^{(w)} \nabla \psi_j) \cdot (\sum_k c_k^{(w)} \nabla \psi_k), \psi_i).$$
 (275)

It can also be instructive to write the linear systems using matrices and ectors. Define K as the matrix corresponding to the Laplace operator ∇^2 . That, $K_{i,j} = (\nabla \psi_j, \nabla \psi_i)$. Let us introduce the vectors

$$b^{(w)} = (b_0^{(w)}, \dots, b_N^{(w)}),$$

$$b^{(T)} = (b_0^{(T)}, \dots, b_N^{(T)}),$$

$$c^{(w)} = (c_0^{(w)}, \dots, c_N^{(w)}),$$

$$c^{(T)} = (c_0^{(T)}, \dots, c_N^{(T)}).$$

he system (270)-(271) can now be expressed in matrix-vector form as

$$\mu K c^{(w)} = b^{(w)}, \tag{276}$$

$$\kappa K c^{(T)} = b^{(T)} \,. \tag{277}$$

We can solve the first system for $c^{(w)}$, and then the right-hand side $b^{(T)}$ is nown such that we can solve the second system for $c^{(T)}$.

Coupled linear systems. Despite the fact that w can be computed first, ithout knowing T, we shall now pretend that w and T enter a two-way coupling 1ch that we need to derive the algebraic equations as one system for all the 1ch nknowns $c_j^{(w)}$ and $c_j^{(T)}$, $j=0,\ldots,N$. This system is nonlinear in $c_j^{(w)}$ because f the $\nabla w \cdot \nabla w$ product. To remove this nonlinearity, imagine that we introduce 1ch iteration method where we replace $\nabla w \cdot \nabla w$ by $\nabla w_- \cdot \nabla w$, w_- being the w 1ch previous iteration. Then the term $\nabla w_- \cdot \nabla w$ is linear in w 1ch nce w_- is known. The total linear system becomes

$$\begin{split} \sum_{j=0}^{N} A_{i,j}^{(w,w)} c_{j}^{(w)} + \sum_{j=0}^{N} A_{i,j}^{(w,T)} c_{j}^{(T)} &= b_{i}^{(w)}, \quad i = 0, \dots, N, \\ \sum_{j=0}^{N} A_{i,j}^{(T,w)} c_{j}^{(w)} + \sum_{j=0}^{N} A_{i,j}^{(T,T)} c_{j}^{(T)} &= b_{i}^{(T)}, \quad i = 0, \dots, N, \\ A_{i,j}^{(w,w)} &= \mu(\nabla \psi_{j}, \psi_{i}), \\ A_{i,j}^{(w,T)} &= 0, \\ b_{i}^{(w)} &= (\beta, \psi_{i}), \\ A_{i,j}^{(w,T)} &= \mu((\nabla \psi w_{-}) \cdot \nabla \psi_{j}), \psi_{i}), \\ A_{i,j}^{(T,T)} &= \kappa(\nabla \psi_{j}, \psi_{i}), \\ b_{i}^{(T)} &= 0. \end{split}$$

This system can alternatively be written in matrix-vector form as

$$\mu K c^{(w)} = 0 b^{(w)},$$

 $L c^{(w)} + \kappa K c^{(T)} = 0.$

with L as the matrix from the $\nabla w_- \cdot \nabla$ operator: $L_{i,j} = A_{i,j}^{(w,T)}$. The matrix-vector equations are often conveniently written in block

$$\left(\begin{array}{cc} \mu K & 0 \\ L & \kappa K \end{array} \right) \left(\begin{array}{c} c^{(w)} \\ c^{(T)} \end{array} \right) = \left(\begin{array}{c} b^{(w)} \\ 0 \end{array} \right),$$

Note that in the general case where all unknowns enter all equati have to solve the compound system (297)-(298) since then we cannot ut special property that (270) does not involve T and can be solved first.

When the viscosity depends on the temperature, the $\mu \nabla^2 w$ term replaced by $\nabla \cdot (\mu(T)\nabla w)$, and then T enters the equation for w. Now a two-way coupling since both equations contain w and T and therefo be solved simultaneously Th equation $\nabla \cdot (\mu(T)\nabla w) = -\beta$ is nonlinear some iteration procedure is invoked, where we use a previously compute the viscosity $(\mu(T_-))$, the coefficient is known, and the equation involved one unknown, w. In that case we are back to the one-way coupled set of

We may also formulate our PDE system as a vector equation. To this introduce the vector of unknowns $\boldsymbol{u}=(u^{(0)},u^{(1)})$, where $u^{(0)}=w$ and v We then have

$$\nabla^2 \boldsymbol{u} = \left(\begin{array}{c} -\mu^{-1}\beta \\ -\kappa^{-1}\mu \nabla u^{(0)} \cdot \nabla u^{(0)} \end{array} \right).$$

0.4 Different function spaces for the unknowns

is easy to generalize the previous formulation to the case where $w \in V^{(w)}$ and $V^{(T)}$, where $V^{(w)}$ and $V^{(T)}$ can be different spaces with different numbers f degrees of freedom. For example, we may use quadratic basis functions for and linear for T. Approximation of the unknowns by different finite element paces is known as *mixed finite element methods*.

We write

$$V^{(w)} = \operatorname{span}\{\psi_0^{(w)}, \dots, \psi_{N_w}^{(w)}\},\$$
$$V^{(T)} = \operatorname{span}\{\psi_0^{(T)}, \dots, \psi_{N_T}^{(T)}\}.$$

he next step is to multiply (260) by a test function $v^{(w)} \in V^{(w)}$ and (261) by a $v^{(T)} \in V^{(T)}$, integrate by parts and arrive at

$$\int_{\Omega} \mu \nabla w \cdot \nabla v^{(w)} \, \mathrm{d}x = \int_{\Omega} \beta v^{(w)} \, \mathrm{d}x \quad \forall v^{(w)} \in V^{(w)}, \tag{288}$$

$$\int_{\Omega} \kappa \nabla T \cdot \nabla v^{(T)} \, \mathrm{d}x = \int_{\Omega} \mu \nabla w \cdot \nabla w \, v^{(T)} \, \mathrm{d}x \quad \forall v^{(T)} \in V^{(T)}. \tag{289}$$

The compound scalar variational formulation applies a test vector function $= (v^{(w)}, v^{(T)})$ and reads

$$\int_{\Omega} (\mu \nabla w \cdot \nabla v^{(w)} + \kappa \nabla T \cdot \nabla v^{(T)}) \, \mathrm{d}x = \int_{\Omega} (\beta v^{(w)} + \mu \nabla w \cdot \nabla w \, v^{(T)}) \, \mathrm{d}x, \quad (290)$$

alid $\forall \boldsymbol{v} \in \boldsymbol{V} = V^{(w)} \times V^{(T)}$.

The associated linear system is similar to (270)-(271) or (297)-(298), except nat we need to distinguish between $\psi_i^{(w)}$ and $\psi_i^{(T)}$, and the range in the sums ver j must match the number of degrees of freedom in the spaces $V^{(w)}$ and $\dot{V}^{(T)}$. The formulas become

$$\sum_{j=0}^{N_w} A_{i,j}^{(w)} c_j^{(w)} = b_i^{(w)}, \quad i = 0, \dots, N_w,$$
(291)

$$\sum_{j=0}^{N_T} A_{i,j}^{(T)} c_j^{(T)} = b_i^{(T)}, \quad i = 0, \dots, N_T,$$
(292)

$$A_{i,j}^{(w)} = \mu(\nabla \psi_j^{(w)}, \psi_i^{(w)}), \tag{293}$$

$$b_i^{(w)} = (\beta, \psi_i^{(w)}),$$
 (294)

$$A_{i,j}^{(T)} = \kappa(\nabla \psi_j^{(T)}, \psi_i^{(T)}), \tag{295}$$

$$b_i^{(T)} = \mu(\nabla w_-, \psi_i^{(T)}). \tag{296}$$

In the case we formulate one compound linear system involving by $j = 0, ..., N_w$, and $c_i^{(T)}$, $j = 0, ..., N_T$, (297)-(298) becomes

$$\sum_{j=0}^{N_w} A_{i,j}^{(w,w)} c_j^{(w)} + \sum_{j=0}^{N_T} A_{i,j}^{(w,T)} c_j^{(T)} = b_i^{(w)}, \quad i = 0, \dots, N_w,$$

$$\sum_{j=0}^{N_w} A_{i,j}^{(T,w)} c_j^{(w)} + \sum_{j=0}^{N_T} A_{i,j}^{(T,T)} c_j^{(T)} = b_i^{(T)}, \quad i = 0, \dots, N_T,$$

$$A_{i,j}^{(w,w)} = \mu(\nabla \psi_j^{(w)}, \psi_i^{(w)}),$$

$$A_{i,j}^{(w,T)} = 0,$$

$$b_i^{(w)} = (\beta, \psi_i^{(w)}),$$

$$A_{i,j}^{(w,T)} = \mu(\nabla w_- \cdot \nabla \psi_j^{(w)}), \psi_i^{(T)}),$$

$$A_{i,j}^{(T,T)} = \kappa(\nabla \psi_j^{(T)}, \psi_i^{(T)}),$$

$$b_i^{(T)} = 0.$$

The corresponding block form

$$\left(\begin{array}{cc} \mu K^{(w)} & 0 \\ L & \kappa K^{(T)} \end{array} \right) \left(\begin{array}{c} c^{(w)} \\ c^{(T)} \end{array} \right) = \left(\begin{array}{c} b^{(w)} \\ 0 \end{array} \right),$$

has square and rectangular block matrices: $K^{(w)}$ is $N_w \times N_w$, $K^{(T)}$ is N_w while L is $N_T \times N_w$,

20.5 Computations in 1D

We can reduce the system (260)-(261) to one space dimension, which corr to flow in a channel between two flat plates. Alternatively, one may a flow in a circular pipe, introduce cylindrical coordinates, and utilize the symmetry to reduce the equations to a one-dimensional problem in the coordinate. The former model becomes

$$\mu w_{xx} = -\beta,$$

$$\kappa T_{xx} = -\mu w_x^2.$$

while the model in the radial coordinate r reads

$$\mu \frac{1}{r} \frac{d}{dr} \left(r \frac{dw}{dr} \right) = -\beta,$$

$$\kappa \frac{1}{r} \frac{d}{dr} \left(r \frac{dT}{dr} \right) = -\mu \left(\frac{dw}{dr} \right)^2.$$

The domain for (305)-(306) is $\Omega = [0, H]$, with boundary conditions w(0) = v(H) = 0 and $T(0) = T(H) = T_0$. For (307)-(308) the domain is [0, R] (R being ne radius of the pipe) and the boundary conditions are du/dr = dT/dr = 0 for t=0, t=0, and t=0, and t=0.

Calculations to be continued...

1 Exercises

exercise 23: Refactor functions into a more general class

ection 11.2 displays three functions for computing the analytical solution of ome simple model problems. There is quite some repetitive code, suggesting at the functions can benefit from being refactored into a class where the user an define the f(x), a(x), and the boundary conditions in particular methods a subclasses. Demonstrate how the new class can be used to solve the three articular problems in Section 11.2.

In the method that computes the solution, check that the solution found fulfills ne differential equation and the boundary conditions. Filename: $uxx_f_sympy_class$

exercise 24: Compute the deflection of a cable with sine unctions

hanging cable of length L with significant tension has a downward deflection $\cdot(x)$ governed by

Solve

$$Tw''(x) = \ell(x),$$

here T is the tension in the cable and $\ell(x)$ the load per unit length. The cable fixed at x=0 and x=L so the boundary conditions become T(0)=T(L)=0. We assume a constant load $\ell(x)=\mathrm{const.}$

The solution is expected to be symmetric around x=L/2. Formulating the roblem for $x \in \Omega = [0, L/2]$ and then scaling it, results in the scaled problem or the dimensionless vertical deflection u:

$$u'' = 1$$
, $x \in (0,1)$, $u(0) = 0$, $u'(1) = 0$.

it roduce the function space spanned by $\psi_i = \sin((i+1)\pi x/2), i=1,\ldots,N$. se a Galerkin and a least squares method to find the coefficients c_j in $u(x) = \sum_j c_j \psi_j$. Find how fast the coefficients decrease in magnitude by looking at i/c_{j-1} . Find the error in the maximum deflection at x=1 when only one basis motion is used (N=0).

What happens if we choose basis functions $\psi_i = \sin((i+1)\pi x)$? These basis unctions are appropriate if we do not utilize symmetry and solve the problem n [0, L]. A scaled version of this problem reads

$$u'' = 1$$
, $x \in (0,1)$, $u(0) = u(1) = 0$.

Carry out the computations with N=0 and demonstrate that the madeflection u(1/2) is the same in the problem utilizing symmetry and the provening the whole cable. Filename: cable_sin.pdf.

Exercise 25: Check integration by parts

Consider the Galerkin method for the problem involving u in Exercise 2 that the formulas for c_j are independent of whether we perform integraparts or not. Filename: cable_integr_by_parts.pdf.

Exercise 26: Compute the deflection of a cable with elements

Solve the problem for u in Exercise 24 using two P1 linear elements. Findable_2P1.pdf.

Exercise 27: Compute the deflection of a cable with element

Solve the problem for u in Exercise 24 using one P2 element with quadratunctions. Filename: cable 1P2.pdf.

Exercise 28: Compute the deflection of a cable with a load

We consider the deflection of a tension cable as described in Exercise 2 the load is

$$\ell(x) = \begin{cases} \ell_1, & x < L/2, \\ \ell_2, & x \ge L/2 \end{cases} \quad x \in [0, L].$$

This load is not symmetric with respect to the midpoint x = L/2 so the loses its symmetry and we must solve the scaled problem

$$u'' = \begin{cases} 1, & x < 1/2, \\ 0, & x \ge 1/2 \end{cases}$$
 $x \in (0,1), u(0) = 0, u(1) = 0.$

- a) Use $\psi_i = \sin((i+1)\pi x)$, $i=0,\ldots,N$ and the Galerkin method integration by parts. Derive a formula for c_j in the solution expansi $\sum_j c_j \psi_j$. Plot how fast the coefficients c_j tend to zero (on a log scale).
- b) Solve the problem with P1 finite elements. Plot the solution for N_e elements.

Filename: cable discont load.pdf.

exercise 29: Show equivalence between linear systems

icorporation of Dirichlet conditions at x=0 and x=L in a finite element tesh on $\Omega=[0,L]$ can either be done by introducing an expansion $u(x)=c_0\varphi_0+U_{N_n}\varphi_{N_n}+\sum_{j=0}^N c_j\varphi_{\nu(j)}$, with $N=N_n-2$ and considering u values to the inner nodes as unknowns, or one can assemble the matrix system with $(x)=\sum_{j=0}^{N=N_n} c_j\varphi_j$ and afterwards replace the rows corresponding to known c_j alues by the boundary conditions. Show that the two approaches are equivalent.

exercise 30: Compute with a non-uniform mesh

verive the linear system for the problem -u'' = 2 on [0,1], with u(0) = 0 and u(1) = 1, using P1 elements and a *non-uniform* mesh. The vertices have pordinates $x_0 = 0 < x_1 < \cdots < x_N = 1$, and the length of cell number e is $e = x_{e+1} - x_e$.

It is of interest to compare the discrete equations for the finite element iethod in a non-uniform mesh with the corresponding discrete equations arising om a finite difference method. Go through the derivation of the finite difference ormula $u''(x_i) \approx [D_x D_x u]_i$ and modify it to find a natural discretization of $u''(x_i)$ on a non-uniform mesh. Filename: nonuniform_P1.pdf.

'roblem 31: Solve a 1D finite element problem by hand

he following scaled 1D problem is a very simple, yet relevant, model for provective transport in fluids:

$$u' = \epsilon u'', \quad u(0) = 0, \ u(1) = 1, \ x \in [0, 1].$$
 (309)

-) Find the analytical solution to this problem. (Introduce w=u', solve the rst-order differential equation for w(x), and integrate once more.)
-) Derive the variational form of this problem.
-) Introduce a finite element mesh with uniform partitioning. Use P1 elements and compute the element matrix and vector for a general element.
-) Incorporate the boundary conditions and assemble the element contributions.
-) Identify the resulting linear system as a finite difference discretization of the ifferential equation using

$$[D_{2x}u = \epsilon D_x D_x u]_i.$$

Compute the numerical solution and plot it together with the exact solution or a mesh with 20 elements and $\epsilon=10,1,0.1,0.01$. ilename: convdiff1D_P1.pdf.

Exercise 32: Compare finite elements and differences radially symmetric Poisson equation

We consider the Poisson problem in a disk with radius R with Dirichlet co at the boundary. Given that the solution is radially symmetric and dependent only on the radial coordinate $(r = \sqrt{x^2 + y^2})$, we can red problem to a 1D Poisson equation

$$-\frac{1}{r}\frac{d}{dr}\left(r\frac{du}{dr}\right) = f(r), \quad r \in (0, R), \ u'(0) = 0, \ u(R) = U_R.$$

- a) Derive a variational form of (310) by integrating over the whole posed equivalently: use a weighting function $2\pi rv(r)$ and integrate r fr R.
- b) Use a uniform mesh partition with P1 elements and show what the r set of equations becomes. Integrate the matrix entries exact by hand, by Trapezoidal rule to integrate the f term.
- c) Explain that an intuitive finite difference method applied to (310) g

$$\frac{1}{r_i} \frac{1}{h^2} \left(r_{i+\frac{1}{2}} (u_{i+1} - u_i) - r_{i-\frac{1}{2}} (u_i - u_{i-1}) \right) = f_i, \quad i = rh.$$

For i = 0 the factor $1/r_i$ seemingly becomes problematic. One must have u'(0) = 0, because of the radial symmetry, which implies $u_{-1} = i$ allow introduction of a fictitious value u_{-1} . Using this u_{-1} in the di equation for i = 0 gives

$$\begin{split} \frac{1}{r_0} \frac{1}{h^2} \left(r_{\frac{1}{2}}(u_1 - u_0) - r_{-\frac{1}{2}}(u_0 - u_1) \right) &= \\ \frac{1}{r_0} \frac{1}{2h^2} \left((r_0 + r_1)(u_1 - u_0) - (r_{-1} + r_0)(u_0 - u_1) \right) &\approx 2(u_1 - u_0) \end{split}$$

if we use $r_{-1} + r_1 \approx 2r_0$.

Set up the complete set of equations for the finite difference method compare to the finite element method in case a Trapezoidal rule is integrate the f term in the latter method.

Filename: radial_Poisson1D_P1.pdf.

Exercise 33: Compute with variable coefficients and ements by hand

Consider the problem

$$-\frac{d}{dx}\left(a(x)\frac{du}{dx}\right) + \gamma u = f(x), \quad x \in \Omega = [0, L], \quad u(0) = \alpha, \ u'(L) = \beta$$

We choose $a(x) = 1 + x^2$. Then

$$u(x) = \alpha + \beta(1 + L^2) \tan^{-1}(x), \tag{312}$$

an exact solution if $f(x) = \gamma u$.

Derive a variational formulation and compute general expressions for the lement matrix and vector in an arbitrary element, using P1 elements and a niform partitioning of [0, L]. The right-hand side integral is challenging and an be computed by a numerical integration rule. The Trapezoidal rule (101) ives particularly simple expressions. Filename: $atan1D_P1.pdf$.

exercise 34: Solve a 2D Poisson equation using polynomials nd sines

he classical problem of applying a torque to the ends of a rod can be modeled y a Poisson equation defined in the cross section Ω :

$$-\nabla^2 u = 2, \quad (x, y) \in \Omega,$$

ith u=0 on $\partial\Omega$. Exactly the same problem arises for the deflection of a numbrane with shape Ω under a constant load.

For a circular cross section one can readily find an analytical solution. For a ectangular cross section the analytical approach ends up with a sine series. The lea in this exercise is to use a single basis function to obtain an approximate nswer.

We assume for simplicity that the cross section is the unit square: $\Omega = 0, 1] \times [0, 1]$.

-) We consider the basis $\psi_{p,q}(x,y) = \sin((p+1)\pi x)\sin(q\pi y), p,q=0,\ldots,n$. hese basis functions fulfill the Dirichlet condition. Use a Galerkin method and =0.
-) The basis function involving sine functions are orthogonal. Use this property in the Galerkin method to derive the coefficients $c_{p,q}$ in a formula $u = \sum_{p} \sum_{q} c_{p,q} \psi_{p,q}(x,y)$.
-) Another possible basis is $\psi_i(x,y) = (x(1-x)y(1-y))^{i+1}$, $i=0,\ldots,N$. Use ne Galerkin method to compute the solution for N=0. Which choice of a ngle basis function is best, $u \sim x(1-x)y(1-y)$ or $u \sim \sin(\pi x)\sin(\pi y)$? In rder to answer the question, it is necessary to search the web or the literature or an accurate estimate of the maximum u value at x=y=1/2. ilename: torsion $\sin xy.pdf$.

Exercise 35: Analyze a Crank-Nicolson scheme for the difusion equation

erform the analysis in Section 19.10 for a 1D diffusion equation $u_t = \alpha u_{xx}$ iscretized by the Crank-Nicolson scheme in time:

$$\frac{u^{n+1} - u^n}{\Delta t} = \alpha \frac{1}{2} \left(\frac{\partial u^{n+1}}{\partial x^2} \frac{\partial u^n}{\partial x^2} \right),$$

or written compactly with finite difference operators,

$$[D_t u = \alpha D_x D_x \overline{u}^t]^{n + \frac{1}{2}}.$$

(From a strict mathematical point of view, the u^n and u^{n+1} in these edshould be replaced by u_e^n and u_e^{n+1} to indicate that the unknown is the solution of the PDE discretized in time, but not yet in space, see Section Make plots similar to those in Section 19.10. Filename: fe_diffusion

References

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